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(54) Title: COMPOUNDS FOR IMMUNOTHERAPY AND DIAGNOSIS OF BREAST CANCER AND METHODS FOR THEIR USE

(57) Abstract

Compounds and methods for the treatment and diagnosis of breast cancer are provided. The inventive compounds include polypeptides containing at least a portion of a breast tumor protein. Vaccines and pharmaceutical compositions for immunotherapy of breast cancer comprising such polypeptides, or polynucleotides encoding such polypeptides, are also provided, together with polynucleotides for preparing the inventive polypeptides.

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**COMPOUNDS FOR IMMUNOTHERAPY AND DIAGNOSIS
OF BREAST CANCER AND METHODS FOR THEIR USE**

5 TECHNICAL FIELD

The present invention relates generally to compositions and methods for the treatment and diagnosis of breast cancer. The invention is more particularly related to polypeptides comprising at least a portion of a protein that is preferentially expressed in breast tumor tissue and to polynucleotides encoding such polypeptides.

10 Such polypeptides and polynucleotides may be used in vaccines and pharmaceutical compositions for treatment of breast cancer. Additionally such polypeptides and polynucleotides may be used in the immunodiagnosis of breast cancer.

BACKGROUND OF THE INVENTION

15 Breast cancer is a significant health problem for women in the United States and throughout the world. Although advances have been made in detection and treatment of the disease, breast cancer remains the second leading cause of cancer-related deaths in women, affecting more than 180,000 women in the United States each year. For women in North America, the life-time odds of getting breast

20 cancer are now one in eight.

No vaccine or other universally successful method for the prevention or treatment of breast cancer is currently available. Management of the disease currently relies on a combination of early diagnosis (through routine breast screening procedures) and aggressive treatment, which may include one or more of a variety of

25 treatments such as surgery, radiotherapy, chemotherapy and hormone therapy. The course of treatment for a particular breast cancer is often selected based on a variety of prognostic parameters, including an analysis of specific tumor markers. See, e.g., Porter-Jordan and Lippman, *Breast Cancer* 8:73-100 (1994). However, the use of established markers often leads to a result that is difficult to interpret, and the high

mortality observed in breast cancer patients indicates that improvements are needed in the treatment, diagnosis and prevention of the disease.

Accordingly, there is a need in the art for improved methods for therapy and diagnosis of breast cancer. The present invention fulfills these needs and
5 further provides other related advantages.

SUMMARY OF THE INVENTION

The present invention provides compounds and methods for immunotherapy of breast cancer. In one aspect, isolated polypeptides are provided
10 comprising at least an immunogenic portion of a breast tumor protein or a variant of said protein that differs only in conservative substitutions and/or modifications, wherein the breast tumor protein comprises an amino acid sequence encoded by a polynucleotide comprising a sequence selected from the group consisting of (a) nucleotide sequences recited in SEQ ID NOS: 3, 10, 17, 24, 45-52, 55-67, 72, 73, 89-
15 97, 102 and 107, (b) complements of said nucleotide sequences and (c) sequences that hybridize to a sequence of (a) or (b) under moderately stringent conditions. In specific embodiments, the isolated polypeptides of the present invention comprise an amino acid sequence of SEQ ID NO: 98, 99 or 101.

In related aspects, isolated polynucleotides encoding the above
20 polypeptides are provided. In specific embodiments, such polynucleotides comprise sequences provided in SEQ ID NOS: 3, 10, 17, 24, 45-52 and 55-67, 72, 73, 89-97, 102 and 107. The present invention further provides expression vectors comprising the above polynucleotides and host cells transformed or transfected with such expression vectors. In preferred embodiments, the host cells are selected from the
25 group consisting of *E. coli*, yeast and mammalian cells.

In another aspect, the present invention provides fusion proteins comprising a first and a second inventive polypeptide or, alternatively, an inventive polypeptide and a known breast antigen.

The present invention also provides pharmaceutical compositions
30 comprising at least one of the above polypeptides, or a polynucleotide encoding such a polypeptide, and a physiologically acceptable carrier, together with vaccines

comprising at least one or more such polypeptide or polynucleotide in combination with a non-specific immune response enhancer. Pharmaceutical compositions and vaccines comprising one or more of the above fusion proteins are also provided.

In related aspects, pharmaceutical compositions for the treatment of 5 breast cancer comprising at least one polypeptide and a physiologically acceptable carrier are provided, wherein the polypeptide comprises an immunogenic portion of a breast tumor protein or a variant thereof, the breast tumor protein being encoded by a polynucleotide comprising a sequence selected from the group consisting of: (a) nucleotide sequences recited in SEQ ID NOS: 1, 2, 4-9, 11-16, 18-23, 25-44, 53, 54, 10 68-71, 74-88 and 103-106, (b) complements of said nucleotide sequences, and (c) sequences that hybridize to a sequence of (a) or (b) under moderately stringent conditions. The invention also provides vaccines for the treatment of breast cancer comprising such polypeptides in combination with a non-specific immune response enhancer, together with pharmaceutical compositions and vaccines comprising at 15 least one polynucleotide comprising a sequence provided in SEQ ID NOS: 1, 2, 4-9, 11-16, 18-23, 25-44, 53, 54, 68-71, 74-88 and 103-106.

In yet another aspect, methods are provided for inhibiting the development of breast cancer in a patient, comprising administering an effective amount of at least one of the above pharmaceutical compositions and/or vaccines.

20 The present invention also provides methods for immunodiagnosis of breast cancer, together with kits for use in such methods. In one specific aspect of the present invention, methods are provided for detecting breast cancer in a patient, comprising: (a) contacting a biological sample obtained from a patient with a binding agent that is capable of binding to one of the above polypeptides; and (b) detecting in 25 the sample a protein or polypeptide that binds to the binding agent. In preferred embodiments, the binding agent is an antibody, most preferably a monoclonal antibody.

In related aspects, methods are provided for monitoring the progression of breast cancer in a patient, comprising: (a) contacting a biological 30 sample obtained from a patient with a binding agent that is capable of binding to one

of the above polypeptides; (b) determining in the sample an amount of a protein or polypeptide that binds to the binding agent; (c) repeating steps (a) and (b); and comparing the amounts of polypeptide detected in steps (b) and (c).

Within related aspects, the present invention provides antibodies, 5 preferably monoclonal antibodies, that bind to the inventive polypeptides, as well as diagnostic kits comprising such antibodies, and methods of using such antibodies to inhibit the development of breast cancer.

The present invention further provides methods for detecting breast cancer comprising: (a) obtaining a biological sample from a patient; (b) contacting 10 the sample with a first and a second oligonucleotide primer in a polymerase chain reaction, at least one of the oligonucleotide primers being specific for a polynucleotide that encodes one of the above polypeptides; and (c) detecting in the sample a DNA sequence that amplifies in the presence of the first and second oligonucleotide primers. In a preferred embodiment, at least one of the 15 oligonucleotide primers comprises at least about 10 contiguous nucleotides of a polynucleotide comprising a sequence selected from the group consisting of SEQ ID NOS: 1-97, 100 and 102-107.

In a further aspect, the present invention provides a method for detecting breast cancer in a patient comprising: (a) obtaining a biological sample 20 from the patient; (b) contacting the sample with an oligonucleotide probe specific for a polynucleotide that encodes one of the above polypeptides; and (c) detecting in the sample a polynucleotide sequence that hybridizes to the oligonucleotide probe. Preferably, the oligonucleotide probe comprises at least about 15 contiguous nucleotides of a polynucleotide comprising a sequence selected from the group 25 consisting of SEQ ID NOS: 1-97, 100 and 102-107.

In related aspects, diagnostic kits comprising the above oligonucleotide probes or primers are provided.

These and other aspects of the present invention will become apparent upon reference to the following detailed description. All references disclosed herein

are hereby incorporated by reference in their entirety as if each was incorporated individually.

5 BRIEF DESCRIPTION OF THE DRAWINGS AND SEQUENCE IDENTIFIERS

Figs. 1A and B show the specific lytic activity of a first and a second B511S-specific CTL clone, respectively, measured on autologous LCL transduced with B511s (filled squares) or HLA-A3 (open squares).

- 10 SEQ ID NO: 1 is the determined 3'cDNA sequence of 1T-5120
SEQ ID NO: 2 is the determined 3'cDNA sequence of 1T-5122
SEQ ID NO: 3 is the determined 3'cDNA sequence of 1T-5123
SEQ ID NO: 4 is the determined 3'cDNA sequence of 1T-5125
SEQ ID NO: 5 is the determined 3'cDNA sequence of 1T-5126
- 15 SEQ ID NO: 6 is the determined 3'cDNA sequence of 1T-5127
SEQ ID NO: 7 is the determined 3'cDNA sequence of 1T-5129
SEQ ID NO: 8 is the determined 3'cDNA sequence of 1T-5130
SEQ ID NO: 9 is the determined 3'cDNA sequence of 1T-5133
SEQ ID NO: 10 is the determined 3'cDNA sequence of 1T-5136
- 20 SEQ ID NO: 11 is the determined 3'cDNA sequence of 1T-5137
SEQ ID NO: 12 is the determined 3'cDNA sequence of 1T-5139
SEQ ID NO: 13 is the determined 3'cDNA sequence of 1T-5142
SEQ ID NO: 14 is the determined 3'cDNA sequence of 1T-5143
SEQ ID NO: 15 is the determined 5'cDNA sequence of 1T-5120
- 25 SEQ ID NO: 16 is the determined 5'cDNA sequence of 1T-5122
SEQ ID NO: 17 is the determined 5'cDNA sequence of 1T-5123
SEQ ID NO: 18 is the determined 5'cDNA sequence of 1T-5125
SEQ ID NO: 19 is the determined 5'cDNA sequence of 1T-5126
SEQ ID NO: 20 is the determined 5'cDNA sequence of 1T-5127
- 30 SEQ ID NO: 21 is the determined 5'cDNA sequence of 1T-5129
SEQ ID NO: 22 is the determined 5'cDNA sequence of 1T-5130

- SEQ ID NO: 23 is the determined 5'cDNA sequence of 1T-5133
SEQ ID NO: 24 is the determined 5'cDNA sequence of 1T-5136
SEQ ID NO: 25 is the determined 5'cDNA sequence of 1T-5137
SEQ ID NO: 26 is the determined 5'cDNA sequence of 1T-5139
5 SEQ ID NO: 27 is the determined 5'cDNA sequence of 1T-5142
SEQ ID NO: 28 is the determined 5'cDNA sequence of 1T-5143
SEQ ID NO: 29 is the determined 5'cDNA sequence of 1D-4315
SEQ ID NO: 30 is the determined 5'cDNA sequence of 1D-4311
SEQ ID NO: 31 is the determined 5'cDNA sequence of 1E-4440
10 SEQ ID NO: 32 is the determined 5'cDNA sequence of 1E-4443
SEQ ID NO: 33 is the determined 5'cDNA sequence of 1D-4321
SEQ ID NO: 34 is the determined 5'cDNA sequence of 1D-4310
SEQ ID NO: 35 is the determined 5'cDNA sequence of 1D-4320
SEQ ID NO: 36 is the determined 5'cDNA sequence of 1E-4448
15 SEQ ID NO: 37 is the determined 5'cDNA sequence of 1S-5105
SEQ ID NO: 38 is the determined 5'cDNA sequence of 1S-5110
SEQ ID NO: 39 is the determined 5'cDNA sequence of 1S-5111
SEQ ID NO: 40 is the determined 5'cDNA sequence of 1S-5116
SEQ ID NO: 41 is the determined 5'cDNA sequence of 1S-5114
20 SEQ ID NO: 42 is the determined 5'cDNA sequence of 1S-5115
SEQ ID NO: 43 is the determined 5'cDNA sequence of 1S-5118
SEQ ID NO: 44 is the determined 5'cDNA sequence of 1T-5134
SEQ ID NO: 45 is the determined 5'cDNA sequence of 1E-4441
SEQ ID NO: 46 is the determined 5'cDNA sequence of 1E-4444
25 SEQ ID NO: 47 is the determined 5'cDNA sequence of 1E-4322
SEQ ID NO: 48 is the determined 5'cDNA sequence of 1S-5103
SEQ ID NO: 49 is the determined 5'cDNA sequence of 1S-5107
SEQ ID NO: 50 is the determined 5'cDNA sequence of 1S-5113
SEQ ID NO: 51 is the determined 5'cDNA sequence of 1S-5117
30 SEQ ID NO: 52 is the determined 5'cDNA sequence of 1S-5112

- SEQ ID NO: 53 is the determined cDNA sequence of 1013E11
- SEQ ID NO: 54 is the determined cDNA sequence of 1013H10
- SEQ ID NO: 55 is the determined cDNA sequence of 1017C2
- SEQ ID NO: 56 is the determined cDNA sequence of 1016F8
- 5 SEQ ID NO: 57 is the determined cDNA sequence of 1015F5
- SEQ ID NO: 58 is the determined cDNA sequence of 1017A11
- SEQ ID NO: 59 is the determined cDNA sequence of 1013A11
- SEQ ID NO: 60 is the determined cDNA sequence of 1016D8
- SEQ ID NO: 61 is the determined cDNA sequence of 1016D12
- 10 SEQ ID NO: 62 is the determined cDNA sequence of 1015E8
- SEQ ID NO: 63 is the determined cDNA sequence of 1015D11
- SEQ ID NO: 64 is the determined cDNA sequence of 1012H8
- SEQ ID NO: 65 is the determined cDNA sequence of 1013C8
- SEQ ID NO: 66 is the determined cDNA sequence of 1014B3
- 15 SEQ ID NO: 67 is the determined cDNA sequence of 1015B2
- SEQ ID NO: 68-71 are the determined cDNA sequences of previously identified antigens
- SEQ ID NO: 72 is the determined cDNA sequence of JJ9434
- SEQ ID NO: 73 is the determined cDNA sequence of B535S
- 20 SEQ ID NO: 74-88 are the determined cDNA sequence of previously identified antigens
- SEQ ID NO: 89 is the determined cDNA sequence of B534S
- SEQ ID NO: 90 is the determined cDNA sequence of B538S
- SEQ ID NO: 91 is the determined cDNA sequence of B542S
- 25 SEQ ID NO: 92 is the determined cDNA sequence of B543S
- SEQ ID NO: 93 is the determined cDNA sequence of P501S
- SEQ ID NO: 94 is the determined cDNA sequence of B541S
- SEQ ID NO: 95 is an extended cDNA sequence for 1016F8 (also referred to as B511S)
- 30 SEQ ID NO: 96 is an extended cDNA sequence for 1016D12 (also referred to as

B532S)

SEQ ID NO: 97 is an extended cDNA sequence for 1012H8 (also referred to as B533S)

SEQ ID NO: 98 is the predicted amino acid sequence for B511S

5 SEQ ID NO: 99 is the predicted amino acid sequence for B532S

SEQ ID NO: 100 is the determined full-length cDNA sequence for P501S

SEQ ID NO: 101 is the predicted amino acid sequence for P501S

SEQ ID NO: 102 is the determined cDNA sequence of clone #19605, also referred to as 1017C2, showing no significant homology to any known gene

10 SEQ ID NO: 103 is the determined 3' end cDNA sequence for clone #19599, showing homology to the Tumor Expression Enhanced gene

SEQ ID NO: 104 is the determined 5' end cDNA sequence for clone #19599, showing homology to the Tumor Expression Enhanced gene

SEQ ID NO: 105 is the determined cDNA sequence for clone #19607, showing

15 homology to Stromelysin-3

SEQ ID NO: 106 is the determined cDNA sequence for clone #19601, showing homology to Collagen

SEQ ID NO: 107 is the determined cDNA sequence of clone #19606, also referred to as B546S, showing no significant homology to any known gene

20

DETAILED DESCRIPTION OF THE INVENTION

As noted above, the present invention is generally directed to compositions and methods for the immunotherapy and diagnosis of breast cancer.

The inventive compositions are generally isolated polypeptides that comprise at least

25 a portion of a breast tumor protein. Also included within the present invention are molecules (such as an antibody or fragment thereof) that bind to the inventive polypeptides. Such molecules are referred to herein as "binding agents."

In particular, the subject invention discloses isolated polypeptides comprising at least a portion of a human breast tumor protein, or a variant thereof,

30 wherein the breast tumor protein includes an amino acid sequence encoded by a

polynucleotide molecule including a sequence selected from the group consisting of: nucleotide sequences recited in SEQ ID NOS: 1-97, 100 and 102-107, the complements of said nucleotide sequences, and variants thereof. In certain specific embodiments, the inventive polypeptides comprise an amino acid sequence selected 5 from the group consisting of sequences provided in SEQ ID NO: 98, 99 and 101, and variants thereof. As used herein, the term "polypeptide" encompasses amino acid chains of any length, including full length proteins, wherein the amino acid residues are linked by covalent peptide bonds. Thus, a polypeptide comprising a portion of one of the above breast proteins may consist entirely of the portion, or the portion 10 may be present within a larger polypeptide that contains additional sequences. The additional sequences may be derived from the native protein or may be heterologous, and such sequences may be immunoreactive and/or antigenic.

As used herein, an "immunogenic portion" of a human breast tumor protein is a portion that is capable of eliciting an immune response in a patient 15 inflicted with breast cancer and as such binds to antibodies present within sera from a breast cancer patient. Such immunogenic portions generally comprise at least about 5 amino acid residues, more preferably at least about 10, and most preferably at least about 20 amino acid residues. Immunogenic portions of the proteins described herein may be identified in antibody binding assays. Such assays may generally be 20 performed using any of a variety of means known to those of ordinary skill in the art, as described, for example, in Harlow and Lane, *Antibodies: A Laboratory Manual*, Cold Spring Harbor Laboratory, Cold Spring Harbor, NY, 1988. For example, a polypeptide may be immobilized on a solid support (as described below) and contacted with patient sera to allow binding of antibodies within the sera to the 25 immobilized polypeptide. Unbound sera may then be removed and bound antibodies detected using, for example, ¹²⁵I-labeled Protein A. Alternatively, a polypeptide may be used to generate monoclonal and polyclonal antibodies for use in detection of the polypeptide in blood or other fluids of breast cancer patients. Methods for preparing and identifying immunogenic portions of antigens of known sequence are well known

in the art and include those summarized in Paul, *Fundamental Immunology*, 3rd ed., Raven Press, 1993, pp. 243-247.

The term "polynucleotide(s)," as used herein, means a single or double-stranded polymer of deoxyribonucleotide or ribonucleotide bases and includes

5 DNA and corresponding RNA molecules, including HnRNA and mRNA molecules, both sense and anti-sense strands, and comprehends cDNA, genomic DNA and recombinant DNA, as well as wholly or partially synthesized polynucleotides. An HnRNA molecule contains introns and corresponds to a DNA molecule in a generally one-to-one manner. An mRNA molecule corresponds to an HnRNA and DNA

10 molecule from which the introns have been excised. A polynucleotide may consist of an entire gene, or any portion thereof. Operable anti-sense polynucleotides may comprise a fragment of the corresponding polynucleotide, and the definition of "polynucleotide" therefore includes all such operable anti-sense fragments.

The compositions and methods of the present invention also

15 encompass variants of the above polypeptides and polynucleotides. A polypeptide "variant," as used herein, is a polypeptide that differs from the recited polypeptide only in conservative substitutions and/or modifications, such that the therapeutic, antigenic and/or immunogenic properties of the polypeptide are retained. In a preferred embodiment, variant polypeptides differ from an identified sequence by

20 substitution, deletion or addition of five amino acids or fewer. Such variants may generally be identified by modifying one of the above polypeptide sequences, and evaluating the antigenic properties of the modified polypeptide using, for example, the representative procedures described herein. Polypeptide variants preferably exhibit at least about 70%, more preferably at least about 90% and most preferably at

25 least about 95% identity (determined as described below) to the identified polypeptides.

As used herein, a "conservative substitution" is one in which an amino acid is substituted for another amino acid that has similar properties, such that one skilled in the art of peptide chemistry would expect the secondary structure and

30 hydrophobic nature of the polypeptide to be substantially unchanged. In general, the

following groups of amino acids represent conservative changes: (1) ala, pro, gly, glu, asp, gln, asn, ser, thr; (2) cys, ser, tyr, thr; (3) val, ile, leu, met, ala, phe; (4) lys, arg, his; and (5) phe, tyr, trp, his.

Variants may also, or alternatively, contain other modifications,
5 including the deletion or addition of amino acids that have minimal influence on the antigenic properties, secondary structure and hydrophobic nature of the polypeptide. For example, a polypeptide may be conjugated to a signal (or leader) sequence at the N-terminal end of the protein which co-translationally or post-translationally directs transfer of the protein. The polypeptide may also be conjugated to a linker or other
10 sequence for ease of synthesis, purification or identification of the polypeptide (e.g., poly-His), or to enhance binding of the polypeptide to a solid support. For example, a polypeptide may be conjugated to an immunoglobulin Fc region.

A nucleotide "variant" is a sequence that differs from the recited nucleotide sequence in having one or more nucleotide deletions, substitutions or
15 additions. Such modifications may be readily introduced using standard mutagenesis techniques, such as oligonucleotide-directed site-specific mutagenesis as taught, for example, by Adelman et al. (*DNA*, 2:183, 1983). Nucleotide variants may be naturally occurring allelic variants, or non-naturally occurring variants. Variant nucleotide sequences preferably exhibit at least about 70%, more preferably at least
20 about 80% and most preferably at least about 90% identity (determined as described below) to the recited sequence.

The antigens provided by the present invention include variants that are encoded by DNA sequences which are substantially homologous to one or more of the DNA sequences specifically recited herein. "Substantial homology," as used
25 herein, refers to DNA sequences that are capable of hybridizing under moderately stringent conditions. Suitable moderately stringent conditions include prewashing in a solution of 5X SSC, 0.5% SDS, 1.0 mM EDTA (pH 8.0); hybridizing at 50°C-65° C, 5X SSC, overnight or, in the event of cross-species homology, at 45°C with 0.5X SSC; followed by washing twice at 65°C for 20 minutes with each of 2X, 0.5X and
30 0.2X SSC containing 0.1% SDS. Such hybridizing DNA sequences are also within

the scope of this invention, as are nucleotide sequences that, due to code degeneracy, encode an immunogenic polypeptide that is encoded by a hybridizing DNA sequence.

Two nucleotide or polypeptide sequences are said to be "identical" if the sequence of nucleotides or amino acid residues in the two sequences is the same
5 when aligned for maximum correspondence as described below. Comparisons between two sequences are typically performed by comparing the sequences over a comparison window to identify and compare local regions of sequence similarity. A "comparison window" as used herein, refers to a segment of at least about 20 contiguous positions, usually 30 to about 75, more preferably 40 to about 50, in
10 which a sequence may be compared to a reference sequence of the same number of contiguous positions after the two sequences are optimally aligned.

Optimal alignment of sequences for comparison may be conducted using the Megalign program in the Lasergene suite of bioinformatics software (DNASTAR, Inc., Madison, WI), using default parameters. This program embodies
15 several alignment schemes described in the following references: Dayhoff, M.O. (1978) A model of evolutionary change in proteins – Matrices for detecting distant relationships. In Dayhoff, M.O. (ed.) *Atlas of Protein Sequence and Structure*, National Biomedical Research Foundation, Washington DC Vol. 5, Suppl. 3, pp. 345-358; Hein J. (1990) Unified Approach to Alignment and Phylogenies pp. 626-645
20 *Methods in Enzymology* vol. 183, Academic Press, Inc., San Diego, CA; Higgins, D.G. and Sharp, P.M. (1989) Fast and sensitive multiple sequence alignments on a microcomputer *CABIOS* 5:151-153; Myers, E.W. and Muller W. (1988) Optimal alignments in linear space *CABIOS* 4:11-17; Robinson, E.D. (1971) *Comb. Theor* 11:105; Saitou, N. Nes, M. (1987) The neighbor joining method. A new method for
25 reconstructing phylogenetic trees *Mol. Biol. Evol.* 4:406-425; Sneath, P.H.A. and Sokal, R.R. (1973) *Numerical Taxonomy – the Principles and Practice of Numerical Taxonomy*, Freeman Press, San Francisco, CA; Wilbur, W.J. and Lipman, D.J. (1983) Rapid similarity searches of nucleic acid and protein data banks *Proc. Natl. Acad. Sci. USA* 80:726-730.

Preferably, the "percentage of sequence identity" is determined by comparing two optimally aligned sequences over a window of comparison of at least 20 positions, wherein the portion of the polynucleotide sequence in the comparison window may comprise additions or deletions (i.e. gaps) of 20 percent or less, usually 5 to 15 percent, or 10 to 12 percent, as compared to the reference sequences (which does not comprise additions or deletions) for optimal alignment of the two sequences. The percentage is calculated by determining the number of positions at which the identical nucleic acid bases or amino acid residue occurs in both sequences to yield the number of matched positions, dividing the number of matched positions by the 10 total number of positions in the reference sequence (i.e. the window size) and multiplying the results by 100 to yield the percentage of sequence identity.

Also included in the scope of the present invention are alleles of the genes encoding the nucleotide sequences recited herein. As used herein, an "allele" or "allelic sequence" is an alternative form of the gene which may result from at least 15 one mutation in the nucleic acid sequence. Alleles may result in altered mRNAs or polypeptides whose structure or function may or may not be altered. Any given gene may have none, one, or many allelic forms. Common mutational changes which give rise to alleles are generally ascribed to natural deletions, additions, or substitutions of nucleotides. Each of these types of changes may occur alone or in combination with 20 the others, one or more times in a given sequence.

For breast tumor polypeptides with immunoreactive properties, variants may alternatively be identified by modifying the amino acid sequence of one of the above polypeptides, and evaluating the immunoreactivity of the modified polypeptide. For breast tumor polypeptides useful for the generation of diagnostic binding agents, a 25 variant may be identified by evaluating a modified polypeptide for the ability to generate antibodies that detect the presence or absence of breast cancer. Such modified sequences may be prepared and tested using, for example, the representative procedures described herein.

The breast tumor proteins of the present invention, and polynucleotide 30 molecules encoding such proteins, may be isolated from breast tumor tissue using any

of a variety of methods well known in the art. Polynucleotide sequences corresponding to a gene (or a portion thereof) encoding one of the inventive breast tumor proteins may be isolated from a breast tumor cDNA library using a subtraction technique as described in detail below. Examples of such DNA sequences are 5 provided in SEQ ID NOS: 1- 97, 100 and 102-107. Partial polynucleotide sequences thus obtained may be used to design oligonucleotide primers for the amplification of full-length polynucleotide sequences in a polymerase chain reaction (PCR), using techniques well known in the art (see, for example, Mullis et al., *Cold Spring Harbor Symp. Quant. Biol.*, 51:263, 1987; Erlich ed., *PCR Technology*, Stockton Press, NY, 10 1989). Once a polynucleotide sequence encoding a polypeptide is obtained, any of the above modifications may be readily introduced using standard mutagenesis techniques, such as oligonucleotide-directed site-specific mutagenesis as taught, for example, by Adelman et al. (*DNA*, 2:183, 1983).

The breast tumor polypeptides disclosed herein may also be generated 15 by synthetic or recombinant means. Synthetic polypeptides having fewer than about 100 amino acids, and generally fewer than about 50 amino acids, may be generated using techniques well known to those of ordinary skill in the art. For example, such polypeptides may be synthesized using any of the commercially available solid-phase techniques, such as the Merrifield solid-phase synthesis method, where amino acids 20 are sequentially added to a growing amino acid chain (see, for example, Merrifield, *J. Am. Chem. Soc.* 85:2149-2146, 1963). Equipment for automated synthesis of polypeptides is commercially available from suppliers such as Perkin Elmer/Applied BioSystems Division (Foster City, CA), and may be operated according to the manufacturer's instructions.

25 Alternatively, any of the above polypeptides may be produced recombinantly by inserting a polynucleotide sequence that encodes the polypeptide into an expression vector and expressing the protein in an appropriate host. Any of a variety of expression vectors known to those of ordinary skill in the art may be employed to express recombinant polypeptides of this invention. Expression may be 30 achieved in any appropriate host cell that has been transformed or transfected with an

expression vector containing a polynucleotide molecule that encodes a recombinant polypeptide. Suitable host cells include prokaryotes, yeast and higher eukaryotic cells. Preferably, the host cells employed are *E. coli*, yeast or a mammalian cell line, such as CHO cells. The polynucleotide sequences expressed in this manner may 5 encode naturally occurring polypeptides, portions of naturally occurring polypeptides, or other variants thereof.

In general, regardless of the method of preparation, the polypeptides disclosed herein are prepared in an isolated, substantially pure form (*i.e.*, the polypeptides are homogenous as determined by amino acid composition and primary 10 sequence analysis). Preferably, the polypeptides are at least about 90% pure, more preferably at least about 95% pure and most preferably at least about 99% pure. In certain preferred embodiments, described in more detail below, the substantially pure polypeptides are incorporated into pharmaceutical compositions or vaccines for use in one or more of the methods disclosed herein.

15 In a related aspect, the present invention provides fusion proteins comprising a first and a second inventive polypeptide or, alternatively, a polypeptide of the present invention and a known breast tumor antigen, together with variants of such fusion proteins.

A polynucleotide sequence encoding a fusion protein of the present 20 invention is constructed using known recombinant DNA techniques to assemble separate polynucleotide sequences encoding the first and second polypeptides into an appropriate expression vector. The 3' end of a polynucleotide sequence encoding the first polypeptide is ligated, with or without a peptide linker, to the 5' end of a polynucleotide sequence encoding the second polypeptide so that the reading frames 25 of the sequences are in phase to permit mRNA translation of the two DNA sequences into a single fusion protein that retains the biological activity of both the first and the second polypeptides.

A peptide linker sequence may be employed to separate the first and the second polypeptides by a distance sufficient to ensure that each polypeptide folds 30 into its secondary and tertiary structures. Such a peptide linker sequence is

incorporated into the fusion protein using standard techniques well known in the art. Suitable peptide linker sequences may be chosen based on the following factors: (1) their ability to adopt a flexible extended conformation; (2) their inability to adopt a secondary structure that could interact with functional epitopes on the first and 5 second polypeptides; and (3) the lack of hydrophobic or charged residues that might react with the polypeptide functional epitopes. Preferred peptide linker sequences contain Gly, Asn and Ser residues. Other near neutral amino acids, such as Thr and Ala may also be used in the linker sequence. Amino acid sequences which may be usefully employed as linkers include those disclosed in Maratea et al., *Gene* 40:39-46, 10 1985; Murphy et al., *Proc. Natl. Acad. Sci. USA* 83:8258-8262, 1986; U.S. Patent No. 4,935,233 and U.S. Patent No. 4,751,180. The linker sequence may be from 1 to about 50 amino acids in length. Peptide sequences are not required when the first and second polypeptides have non-essential N-terminal amino acid regions that can be used to separate the functional domains and prevent steric interference.

15 The ligated polynucleotide sequences are operably linked to suitable transcriptional or translational regulatory elements. The regulatory elements responsible for expression of polynucleotides are located only 5' to the polynucleotide sequence encoding the first polypeptide. Similarly, stop codons required to end translation and transcription termination signals are only present 3' to the 20 polynucleotide sequence encoding the second polypeptide.

Fusion proteins are also provided that comprise a polypeptide of the present invention together with an unrelated immunogenic protein. Preferably the immunogenic protein is capable of eliciting a recall response. Examples of such proteins include tetanus, tuberculosis and hepatitis proteins (see, for example, Stoute 25 et al. *New Engl. J. Med.*, 336:86-91 (1997)).

Polypeptides of the present invention that comprise an immunogenic portion of a breast tumor protein may generally be used for immunotherapy of breast cancer, wherein the polypeptide stimulates the patient's own immune response to breast tumor cells. In further aspects, the present invention provides methods for 30 using one or more of the immunoreactive polypeptides encoded by a polynucleotide

molecule having a sequence provided in SEQ ID NOS: 1- 97, 100 and 102-107 (or fusion proteins comprising one or more such polypeptides and/or polynucleotides encoding such polypeptides) for immunotherapy of breast cancer in a patient. As used herein, a "patient" refers to any warm-blooded animal, preferably a human. A 5 patient may be afflicted with a disease, or may be free of detectable disease. Accordingly, the above immunoreactive polypeptides (or fusion proteins or polynucleotide molecules encoding such polypeptides) may be used to treat breast cancer or to inhibit the development of breast cancer. In a preferred embodiment, the polypeptides are administered either prior to or following surgical removal of primary 10 tumors and/or treatment by administration of radiotherapy and conventional chemotherapeutic drugs.

In these aspects, the polypeptide or fusion protein is generally present within a pharmaceutical composition and/or a vaccine. Pharmaceutical compositions may comprise one or more polypeptides, each of which may contain one or more of 15 the above sequences (or variants thereof), and a physiologically acceptable carrier. The vaccines may comprise one or more of such polypeptides and a non-specific immune response enhancer, wherein the non-specific immune response enhancer is capable of eliciting or enhancing an immune response to an exogenous antigen. Examples of non-specific-immune response enhancers include adjuvants, 20 biodegradable microspheres (e.g., polylactic galactide) and liposomes (into which the polypeptide is incorporated). Pharmaceutical compositions and vaccines may also contain other epitopes of breast tumor antigens, either incorporated into a combination polypeptide (*i.e.*, a single polypeptide that contains multiple epitopes) or present within a separate polypeptide.

25 Alternatively, a pharmaceutical composition or vaccine may contain polynucleotides encoding one or more of the above polypeptides, such that the polypeptide is generated *in situ*. In such pharmaceutical compositions and vaccines, the polynucleotide may be present within any of a variety of delivery systems known to those of ordinary skill in the art, including nucleic acid expression systems, 30 bacteria and viral expression systems. Appropriate nucleic acid expression systems

contain the necessary polynucleotide sequences for expression in the patient (such as a suitable promoter). Bacterial delivery systems involve the administration of a bacterium (such as *Bacillus-Calmette-Guerrin*) that expresses an epitope of a breast tumor cell antigen on its cell surface. In a preferred embodiment, the polynucleotide molecules may be introduced using a viral expression system (e.g., vaccinia or other pox virus, retrovirus, or adenovirus), which may involve the use of a non-pathogenic (defective), replication competent virus. Suitable systems are disclosed, for example, in Fisher-Hoch et al., *PNAS* 86:317-321, 1989; Flexner et al., *Ann. N.Y. Acad. Sci.* 569:86-103, 1989; Flexner et al., *Vaccine* 8:17-21, 1990; U.S. Patent Nos. 4,603,112, 5 4,769,330, and 5,017,487; WO 89/01973; U.S. Patent No. 4,777,127; GB 2,200,651; EP 0,345,242; WO 91/02805; Berkner, *Biotechniques* 6:616-627, 1988; Rosenfeld et 10 al., *Science* 252:431-434, 1991; Kolls et al., *PNAS* 91:215-219, 1994; Kass-Eisler et al., *PNAS* 90:11498-11502, 1993; Guzman et al., *Circulation* 88:2838-2848, 1993; and Guzman et al., *Cir. Res.* 73:1202-1207, 1993. Techniques for incorporating 15 polynucleotides into such expression systems are well known to those of ordinary skill in the art.

The polynucleotides may also be "naked," as described, for example, in published PCT application WO 90/11092, and Ulmer et al., *Science* 259:1745-1749, 1993, reviewed by Cohen, *Science* 259:1691-1692, 1993. The 20 uptake of naked polynucleotides may be increased by coating the polynucleotides onto biodegradable beads, which are efficiently transported into the cells.

Routes and frequency of administration, as well as dosage, will vary from individual to individual and may parallel those currently being used in immunotherapy of other diseases. In general, the pharmaceutical compositions and 25 vaccines may be administered by injection (e.g., intracutaneous, intramuscular, intravenous or subcutaneous), intranasally (e.g., by aspiration) or orally. Between 1 and 10 doses may be administered over a 3-24 week period. Preferably, 4 doses are administered, at an interval of 3 months, and booster administrations may be given periodically thereafter. Alternate protocols may be appropriate for individual 30 patients. A suitable dose is an amount of polypeptide or polynucleotide that is

effective to raise an immune response (cellular and/or humoral) against breast tumor cells in a treated patient. A suitable immune response is at least 10-50% above the basal (*i.e.*, untreated) level. In general, the amount of polypeptide present in a dose (or produced *in situ* by the polynucleotide in a dose) ranges from about 1 pg to about 5 100 mg per kg of host, typically from about 10 pg to about 1 mg, and preferably from about 100 pg to about 1 μ g. Suitable dose sizes will vary with the size of the patient, but will typically range from about 0.01 mL to about 5 mL.

While any suitable carrier known to those of ordinary skill in the art may be employed in the pharmaceutical compositions of this invention, the type of 10 carrier will vary depending on the mode of administration. For parenteral administration, such as subcutaneous injection, the carrier preferably comprises water, saline, alcohol, a lipid, a wax and/or a buffer. For oral administration, any of the above carriers or a solid carrier, such as mannitol, lactose, starch, magnesium stearate, sodium saccharine, talcum, cellulose, glucose, sucrose, and/or magnesium 15 stearate, sodium saccharine, talcum, cellulose, glucose, sucrose, and/or magnesium carbonate, may be employed. Biodegradable microspheres (*e.g.*, polylactic glycolide) may also be employed as carriers for the pharmaceutical compositions of this invention. Suitable biodegradable microspheres are disclosed, for example, in U.S. Patent Nos. 4,897,268 and 5,075,109.

Any of a variety of non-specific immune response enhancers may be 20 employed in the vaccines of this invention. For example, an adjuvant may be included. Most adjuvants contain a substance designed to protect the antigen from rapid catabolism, such as aluminum hydroxide or mineral oil, and a nonspecific stimulator of immune response, such as lipid A, *Bordetella pertussis* or *Mycobacterium tuberculosis*. Such adjuvants are commercially available as, for example, Freund's 25 Incomplete Adjuvant and Complete Adjuvant (Difco Laboratories, Detroit, MI) and Merck Adjuvant 65 (Merck and Company, Inc., Rahway, NJ).

Polypeptides disclosed herein may also be employed in adoptive immunotherapy for the treatment of cancer. Adoptive immunotherapy may be broadly classified into either active or passive immunotherapy. In active 30 immunotherapy, treatment relies on the *in vivo* stimulation of the endogenous host

immune system to react against tumors with the administration of immune response-modifying agents (for example, tumor vaccines, bacterial adjuvants, and/or cytokines).

In passive immunotherapy, treatment involves the delivery of biologic reagents with established tumor-immune reactivity (such as effector cells or antibodies) that can directly or indirectly mediate antitumor effects and does not necessarily depend on an intact host immune system. Examples of effector cells include T lymphocytes (for example, CD8+ cytotoxic T-lymphocyte, CD4+ T-helper, gamma/delta T lymphocytes, tumor-infiltrating lymphocytes), killer cells (such as Natural Killer cells, lymphokine-activated killer cells), B cells, or antigen presenting cells (such as dendritic cells and macrophages) expressing the disclosed antigens. The polypeptides disclosed herein may also be used to generate antibodies or anti-idiotypic antibodies (as in U.S. Patent No. 4,918,164), for passive immunotherapy.

The predominant method of procuring adequate numbers of T-cells for adoptive immunotherapy is to grow immune T-cells *in vitro*. Culture conditions for expanding single antigen-specific T-cells to several billion in number with retention of antigen recognition *in vivo* are well known in the art. These *in vitro* culture conditions typically utilize intermittent stimulation with antigen, often in the presence of cytokines, such as IL-2, and non-dividing feeder cells. As noted above, the immunoreactive polypeptides described herein may be used to rapidly expand antigen-specific T cell cultures in order to generate sufficient number of cells for immunotherapy. In particular, antigen-presenting cells, such as dendritic, macrophage, monocyte, fibroblast or B-cells, may be pulsed with immunoreactive polypeptides or polynucleotide sequence(s) may be introduced into antigen presenting cells, using standard techniques well known in the art. For example, antigen presenting cells may be transfected or transduced with a polynucleotide sequence, wherein said sequence contains a promoter region appropriate for inducing expression, and can be expressed as part of a recombinant virus or other expression system. Several viral vectors may be used to transduce an antigen presenting cell, including pox virus, vaccinia virus, and adenovirus. Antigen presenting cells may be

transfected with polynucleotide sequences disclosed herein by a variety of means, including gene-gun technology, lipid-mediated delivery, electroporation, osmotic shock, and particulate delivery mechanisms, resulting in efficient and acceptable expression levels as determined by one of ordinary skill in the art. For cultured T-
5 cells to be effective in therapy, the cultured T-cells must be able to grow and distribute widely and to survive long term *in vivo*. Studies have demonstrated that cultured T-cells can be induced to grow *in vivo* and to survive long term in substantial numbers by repeated stimulation with antigen supplemented with IL-2 (see, for example, Cheever, M., et al, "Therapy With Cultured T Cells: Principles Revisited,"
10 *Immunological Reviews*, 157:177, 1997).

The polypeptides disclosed herein may also be employed to generate and/or isolate tumor-reactive T-cells, which can then be administered to the patient. In one technique, antigen-specific T-cell lines may be generated by *in vivo* immunization with short peptides corresponding to immunogenic portions of the
15 disclosed polypeptides. The resulting antigen specific CD8+ CTL clones may be isolated from the patient, expanded using standard tissue culture techniques, and returned to the patient.

Alternatively, peptides corresponding to immunogenic portions of the polypeptides may be employed to generate tumor reactive T cell subsets by selective
20 *in vitro* stimulation and expansion of autologous T cells to provide antigen-specific T cells which may be subsequently transferred to the patient as described, for example, by Chang et al. (*Crit. Rev. Oncol. Hematol.*, 22(3), 213, 1996). Cells of the immune system, such as T cells, may be isolated from the peripheral blood of a patient, using a commercially available cell separation system. The separated cells are stimulated
25 with one or more of the immunoreactive polypeptides contained within a delivery vehicle, such as a microsphere, to provide antigen-specific T cells. The population of tumor antigen-specific T cells is then expanded using standard techniques and the cells are administered back to the patient.

In other embodiments, T-cell and/or antibody receptors specific for the
30 polypeptides can be cloned, expanded, and transferred into other vectors or effector

cells for use in adoptive immunotherapy. In particular, T cells may be transfected with the appropriate genes to express the variable domains from tumor specific monoclonal antibodies as the extracellular recognition elements and joined to the T cell receptor signaling chains, resulting in T cell activation, specific lysis, and 5 cytokine release. This enables the T cell to redirect its specificity in an MHC-independent manner. See for example, Eshhar, Z., *Cancer Immunol Immunother*, 45(3-4):131-6, 1997 and Hwu, P., et al, *Cancer Res*, 55(15):3369-73, 1995. Another embodiment may include the transfection of tumor antigen specific alpha and beta T cell receptor chains into alternate T cells, as in Cole, DJ, et al, *Cancer Res*, 55(4):748-10 52, 1995.

In further embodiments, syngeneic or autologous dendritic cells may be pulsed with peptides corresponding to at least an immunogenic portion of a polypeptide disclosed herein. The resulting antigen-specific dendritic cells may either be transferred into a patient, or employed to stimulate T cells to provide 15 antigen-specific T cells which may, in turn, be administered to a patient. The use of peptide-pulsed dendritic cells to generate antigen-specific T cells and the subsequent use of such antigen-specific T cells to eradicate tumors in a murine model has been demonstrated by Cheever et al. (*Immunological Reviews*, 157:177, 1997).

Additionally, vectors expressing the disclosed polynucleotides may be 20 introduced into stem cells taken from the patient and clonally propagated *in vitro* for autologous transplant back into the same patient.

In one specific embodiment, cells of the immune system, such as T cells, may be isolated from the peripheral blood of a patient, using a commercially available cell separation system, such as CellPro Incorporated's (Bothell, WA) 25 CEPRATE™ system (see U.S. Patent No. 5,240,856; U.S. Patent No. 5,215,926; WO 89/06280; WO 91/16116 and WO 92/07243). The separated cells are stimulated with one or more of the immunoreactive polypeptides contained within a delivery vehicle, such as a microsphere, to provide antigen-specific T cells. The population of tumor antigen-specific T cells is then expanded using standard techniques and the cells are 30 administered back to the patient.

Polypeptides of the present invention may also, or alternatively, be used to generate binding agents, such as antibodies or fragments thereof, that are capable of detecting metastatic human breast tumors. Binding agents of the present invention may generally be prepared using methods known to those of ordinary skill 5 in the art, including the representative procedures described herein. Binding agents are capable of differentiating between patients with and without breast cancer, using the representative assays described herein. In other words, antibodies or other binding agents raised against a breast tumor protein, or a suitable portion thereof, will generate a signal indicating the presence of primary or metastatic breast cancer in at 10 least about 20% of patients afflicted with the disease, and will generate a negative signal indicating the absence of the disease in at least about 90% of individuals without primary or metastatic breast cancer. Suitable portions of such breast tumor proteins are portions that are able to generate a binding agent that indicates the presence of primary or metastatic breast cancer in substantially all (*i.e.*, at least about 15 80%, and preferably at least about 90%) of the patients for which breast cancer would be indicated using the full length protein, and that indicate the absence of breast cancer in substantially all of those samples that would be negative when tested with full length protein. The representative assays described below, such as the two-antibody sandwich assay, may generally be employed for evaluating the ability of a 20 binding agent to detect metastatic human breast tumors.

The ability of a polypeptide prepared as described herein to generate antibodies capable of detecting primary or metastatic human breast tumors may generally be evaluated by raising one or more antibodies against the polypeptide (using, for example, a representative method described herein) and determining the 25 ability of such antibodies to detect such tumors in patients. This determination may be made by assaying biological samples from patients with and without primary or metastatic breast cancer for the presence of a polypeptide that binds to the generated antibodies. Such test assays may be performed, for example, using a representative procedure described below. Polypeptides that generate antibodies capable of 30 detecting at least 20% of primary or metastatic breast tumors by such procedures are

considered to be useful in assays for detecting primary or metastatic human breast tumors. Polypeptide specific antibodies may be used alone or in combination to improve sensitivity.

Polypeptides capable of detecting primary or metastatic human breast tumors 5 may be used as markers for diagnosing breast cancer or for monitoring disease progression in patients. In one embodiment, breast cancer in a patient may be diagnosed by evaluating a biological sample obtained from the patient for the level of one or more of the above polypeptides, relative to a predetermined cut-off value. As used herein, suitable "biological samples" include blood, sera and urine.

10 The level of one or more of the above polypeptides may be evaluated using any binding agent specific for the polypeptide(s). A "binding agent," in the context of this invention, is any agent (such as a compound or a cell) that binds to a polypeptide as described above. As used herein, "binding" refers to a noncovalent association between two separate molecules (each of which may be free (*i.e.*, in 15 solution) or present on the surface of a cell or a solid support), such that a "complex" is formed. Such a complex may be free or immobilized (either covalently or noncovalently) on a support material. The ability to bind may generally be evaluated by determining a binding constant for the formation of the complex. The binding constant is the value obtained when the concentration of the complex is divided by 20 the product of the component concentrations. In general, two compounds are said to "bind" in the context of the present invention when the binding constant for complex formation exceeds about 10^3 L/mol. The binding constant may be determined using methods well known to those of ordinary skill in the art.

Any agent that satisfies the above requirements may be a binding 25 agent. For example, a binding agent may be a ribosome with or without a peptide component, an RNA molecule or a peptide. In a preferred embodiment, the binding partner is an antibody, or a fragment thereof. Such antibodies may be polyclonal, or monoclonal. In addition, the antibodies may be single chain, chimeric, CDR-grafted or humanized. Antibodies may be prepared by the methods described herein and by 30 other methods well known to those of skill in the art.

There are a variety of assay formats known to those of ordinary skill in the art for using a binding partner to detect polypeptide markers in a sample. See, e.g., Harlow and Lane, *Antibodies: A Laboratory Manual*, Cold Spring Harbor Laboratory, 1988. In a preferred embodiment, the assay involves the use of binding 5 partner immobilized on a solid support to bind to and remove the polypeptide from the remainder of the sample. The bound polypeptide may then be detected using a second binding partner that contains a reporter group. Suitable second binding partners include antibodies that bind to the binding partner/polypeptide complex. Alternatively, a competitive assay may be utilized, in which a polypeptide is labeled 10 with a reporter group and allowed to bind to the immobilized binding partner after incubation of the binding partner with the sample. The extent to which components of the sample inhibit the binding of the labeled polypeptide to the binding partner is indicative of the reactivity of the sample with the immobilized binding partner.

The solid support may be any material known to those of ordinary skill 15 in the art to which the antigen may be attached. For example, the solid support may be a test well in a microtiter plate or a nitrocellulose or other suitable membrane. Alternatively, the support may be a bead or disc, such as glass, fiberglass, latex or a plastic material such as polystyrene or polyvinylchloride. The support may also be a magnetic particle or a fiber optic sensor, such as those disclosed, for example, in U.S. 20 Patent No. 5,359,681. The binding agent may be immobilized on the solid support using a variety of techniques known to those of skill in the art, which are amply described in the patent and scientific literature. In the context of the present invention, the term "immobilization" refers to both noncovalent association, such as adsorption, and covalent attachment (which may be a direct linkage between the 25 antigen and functional groups on the support or may be a linkage by way of a cross-linking agent). Immobilization by adsorption to a well in a microtiter plate or to a membrane is preferred. In such cases, adsorption may be achieved by contacting the binding agent, in a suitable buffer, with the solid support for a suitable amount of time. The contact time varies with temperature, but is typically between about 1 hour 30 and about 1 day. In general, contacting a well of a plastic microtiter plate (such as

polystyrene or polyvinylchloride) with an amount of binding agent ranging from about 10 ng to about 10 µg, and preferably about 100 ng to about 1 µg, is sufficient to immobilize an adequate amount of binding agent.

Covalent attachment of binding agent to a solid support may generally 5 be achieved by first reacting the support with a bifunctional reagent that will react with both the support and a functional group, such as a hydroxyl or amino group, on the binding agent. For example, the binding agent may be covalently attached to supports having an appropriate polymer coating using benzoquinone or by condensation of an aldehyde group on the support with an amine and an active 10 hydrogen on the binding partner (see, e.g., Pierce Immunotechnology Catalog and Handbook, 1991, at A12-A13).

In certain embodiments, the assay is a two-antibody sandwich assay. This assay may be performed by first contacting an antibody that has been immobilized on a solid support, commonly the well of a microtiter plate, with the 15 sample, such that polypeptides within the sample are allowed to bind to the immobilized antibody. Unbound sample is then removed from the immobilized polypeptide-antibody complexes and a second antibody (containing a reporter group) capable of binding to a different site on the polypeptide is added. The amount of second antibody that remains bound to the solid support is then determined using a 20 method appropriate for the specific reporter group.

More specifically, once the antibody is immobilized on the support as described above, the remaining protein binding sites on the support are typically blocked. Any suitable blocking agent known to those of ordinary skill in the art, such as bovine serum albumin or Tween 20TM (Sigma Chemical Co., St. Louis, MO). The 25 immobilized antibody is then incubated with the sample, and polypeptide is allowed to bind to the antibody. The sample may be diluted with a suitable diluent, such as phosphate-buffered saline (PBS) prior to incubation. In general, an appropriate contact time (*i.e.*, incubation time) is that period of time that is sufficient to detect the presence of polypeptide within a sample obtained from an individual with breast 30 cancer. Preferably, the contact time is sufficient to achieve a level of binding that is

at least about 95% of that achieved at equilibrium between bound and unbound polypeptide. Those of ordinary skill in the art will recognize that the time necessary to achieve equilibrium may be readily determined by assaying the level of binding that occurs over a period of time. At room temperature, an incubation time of about
5 30 minutes is generally sufficient.

Unbound sample may then be removed by washing the solid support with an appropriate buffer, such as PBS containing 0.1% Tween 20TM. The second antibody, which contains a reporter group, may then be added to the solid support. Preferred reporter groups include enzymes (such as horseradish peroxidase),
10 substrates, cofactors, inhibitors, dyes, radionuclides, luminescent groups, fluorescent groups and biotin. The conjugation of antibody to reporter group may be achieved using standard methods known to those of ordinary skill in the art.

The second antibody is then incubated with the immobilized antibody-polypeptide complex for an amount of time sufficient to detect the bound
15 polypeptide. An appropriate amount of time may generally be determined by assaying the level of binding that occurs over a period of time. Unbound second antibody is then removed and bound second antibody is detected using the reporter group. The method employed for detecting the reporter group depends upon the nature of the reporter group. For radioactive groups, scintillation counting or
20 autoradiographic methods are generally appropriate. Spectroscopic methods may be used to detect dyes, luminescent groups and fluorescent groups. Biotin may be detected using avidin, coupled to a different reporter group (commonly a radioactive or fluorescent group or an enzyme). Enzyme reporter groups may generally be detected by the addition of substrate (generally for a specific period of time),
25 followed by spectroscopic or other analysis of the reaction products.

To determine the presence or absence of breast cancer, the signal detected from the reporter group that remains bound to the solid support is generally compared to a signal that corresponds to a predetermined cut-off value. In one preferred embodiment, the cut-off value is the average mean signal obtained when the
30 immobilized antibody is incubated with samples from patients without breast cancer.

In general, a sample generating a signal that is three standard deviations above the predetermined cut-off value is considered positive for breast cancer. In an alternate preferred embodiment, the cut-off value is determined using a Receiver Operator Curve, according to the method of Sackett et al., *Clinical Epidemiology: A Basic Science for Clinical Medicine*, Little Brown and Co., 1985, p. 106-7. Briefly, in this embodiment, the cut-off value may be determined from a plot of pairs of true positive rates (*i.e.*, sensitivity) and false positive rates (100%-specificity) that correspond to each possible cut-off value for the diagnostic test result. The cut-off value on the plot that is the closest to the upper left-hand corner (*i.e.*, the value that encloses the largest area) is the most accurate cut-off value, and a sample generating a signal that is higher than the cut-off value determined by this method may be considered positive. Alternatively, the cut-off value may be shifted to the left along the plot, to minimize the false positive rate, or to the right, to minimize the false negative rate. In general, a sample generating a signal that is higher than the cut-off value determined by this method is considered positive for breast cancer.

In a related embodiment, the assay is performed in a flow-through or strip test format, wherein the antibody is immobilized on a membrane, such as nitrocellulose. In the flow-through test, polypeptides within the sample bind to the immobilized antibody as the sample passes through the membrane. A second, labeled antibody then binds to the antibody-polypeptide complex as a solution containing the second antibody flows through the membrane. The detection of bound second antibody may then be performed as described above. In the strip test format, one end of the membrane to which antibody is bound is immersed in a solution containing the sample. The sample migrates along the membrane through a region containing second antibody and to the area of immobilized antibody. Concentration of second antibody at the area of immobilized antibody indicates the presence of breast cancer. Typically, the concentration of second antibody at that site generates a pattern, such as a line, that can be read visually. The absence of such a pattern indicates a negative result. In general, the amount of antibody immobilized on the membrane is selected to generate a visually discernible pattern when the biological sample contains a level

of polypeptide that would be sufficient to generate a positive signal in the two-antibody sandwich assay, in the format discussed above. Preferably, the amount of antibody immobilized on the membrane ranges from about 25 ng to about 1 µg, and more preferably from about 50 ng to about 500 ng. Such tests can typically be
5 performed with a very small amount of biological sample.

Of course, numerous other assay protocols exist that are suitable for use with the antigens or antibodies of the present invention. The above descriptions are intended to be exemplary only.

In another embodiment, the above polypeptides may be used as
10 markers for the progression of breast cancer. In this embodiment, assays as described above for the diagnosis of breast cancer may be performed over time, and the change in the level of reactive polypeptide(s) evaluated. For example, the assays may be performed every 24-72 hours for a period of 6 months to 1 year, and thereafter performed as needed. In general, breast cancer is progressing in those patients in
15 whom the level of polypeptide detected by the binding agent increases over time. In contrast, breast cancer is not progressing when the level of reactive polypeptide either remains constant or decreases with time.

Antibodies for use in the above methods may be prepared by any of a variety of techniques known to those of ordinary skill in the art. See, e.g., Harlow
20 and Lane, *Antibodies: A Laboratory Manual*, Cold Spring Harbor Laboratory, 1988. In one such technique, an immunogen comprising the antigenic polypeptide is initially injected into any of a wide variety of mammals (e.g., mice, rats, rabbits, sheep and goats). In this step, the polypeptides of this invention may serve as the immunogen without modification. Alternatively, particularly for relatively short
25 polypeptides, a superior immune response may be elicited if the polypeptide is joined to a carrier protein, such as bovine serum albumin or keyhole limpet hemocyanin. The immunogen is injected into the animal host, preferably according to a predetermined schedule incorporating one or more booster immunizations, and the animals are bled periodically. Polyclonal antibodies specific for the polypeptide may

then be purified from such antisera by, for example, affinity chromatography using the polypeptide coupled to a suitable solid support.

Monoclonal antibodies specific for the antigenic polypeptide of interest may be prepared, for example, using the technique of Kohler and Milstein,
5 *Eur. J. Immunol.* 6:511-519, 1976, and improvements thereto. Briefly, these methods involve the preparation of immortal cell lines capable of producing antibodies having the desired specificity (*i.e.*, reactivity with the polypeptide of interest). Such cell lines may be produced, for example, from spleen cells obtained from an animal immunized as described above. The spleen cells are then immortalized by, for
10 example, fusion with a myeloma cell fusion partner, preferably one that is syngeneic with the immunized animal. A variety of fusion techniques may be employed. For example, the spleen cells and myeloma cells may be combined with a nonionic detergent for a few minutes and then plated at low density on a selective medium that supports the growth of hybrid cells, but not myeloma cells. A preferred selection
15 technique uses HAT (hypoxanthine, aminopterin, thymidine) selection. After a sufficient time, usually about 1 to 2 weeks, colonies of hybrids are observed. Single colonies are selected and tested for binding activity against the polypeptide. Hybridomas having high reactivity and specificity are preferred.

Monoclonal antibodies may be isolated from the supernatants of
20 growing hybridoma colonies. In addition, various techniques may be employed to enhance the yield, such as injection of the hybridoma cell line into the peritoneal cavity of a suitable vertebrate host, such as a mouse. Monoclonal antibodies may then be harvested from the ascites fluid or the blood. Contaminants may be removed from the antibodies by conventional techniques, such as chromatography, gel
25 filtration, precipitation, and extraction. The polypeptides of this invention may be used in the purification process in, for example, an affinity chromatography step.

Monoclonal antibodies of the present invention may also be used as therapeutic reagents, to diminish or eliminate breast tumors. The antibodies may be used on their own (for instance, to inhibit metastases) or coupled to one or more
30 therapeutic agents. Suitable agents in this regard include radionuclides,

differentiation inducers, drugs, toxins, and derivatives thereof. Preferred radionuclides include ⁹⁰Y, ¹²³I, ¹²⁵I, ¹³¹I, ¹⁸⁶Re, ¹⁸⁸Re, ²¹¹At, and ²¹²Bi. Preferred drugs include methotrexate, and pyrimidine and purine analogs. Preferred differentiation inducers include phorbol esters and butyric acid. Preferred toxins include ricin, abrin, 5 diphtheria toxin, cholera toxin, gelonin, Pseudomonas exotoxin, Shigella toxin, and pokeweed antiviral protein.

A therapeutic agent may be coupled (*e.g.*, covalently bonded) to a suitable monoclonal antibody either directly or indirectly (*e.g.*, via a linker group). A direct reaction between an agent and an antibody is possible when each possesses a 10 substituent capable of reacting with the other. For example, a nucleophilic group, such as an amino or sulfhydryl group, on one may be capable of reacting with a carbonyl-containing group, such as an anhydride or an acid halide, or with an alkyl group containing a good leaving group (*e.g.*, a halide) on the other.

Alternatively, it may be desirable to couple a therapeutic agent and an 15 antibody via a linker group. A linker group can function as a spacer to distance an antibody from an agent in order to avoid interference with binding capabilities. A linker group can also serve to increase the chemical reactivity of a substituent on an agent or an antibody, and thus increase the coupling efficiency. An increase in chemical reactivity may also facilitate the use of agents, or functional groups on 20 agents, which otherwise would not be possible.

It will be evident to those skilled in the art that a variety of bifunctional or polyfunctional reagents, both homo- and hetero-functional (such as those described in the catalog of the Pierce Chemical Co., Rockford, IL), may be employed as the linker group. Coupling may be effected, for example, through amino 25 groups, carboxyl groups, sulfhydryl groups or oxidized carbohydrate residues. There are numerous references describing such methodology, *e.g.*, U.S. Patent No. 4,671,958, to Rodwell et al.

Where a therapeutic agent is more potent when free from the antibody portion of the immunoconjugates of the present invention, it may be desirable to use a 30 linker group which is cleavable during or upon internalization into a cell. A number

of different cleavable linker groups have been described. The mechanisms for the intracellular release of an agent from these linker groups include cleavage by reduction of a disulfide bond (*e.g.*, U.S. Patent No. 4,489,710, to Spitzer), by irradiation of a photolabile bond (*e.g.*, U.S. Patent No. 4,625,014, to Senter et al.), by hydrolysis of derivatized amino acid side chains (*e.g.*, U.S. Patent No. 4,638,045, to Kohn et al.), by serum complement-mediated hydrolysis (*e.g.*, U.S. Patent No. 4,671,958, to Rodwell et al.), and acid-catalyzed hydrolysis (*e.g.*, U.S. Patent No. 4,569,789, to Blattler et al.).

It may be desirable to couple more than one agent to an antibody. In one embodiment, multiple molecules of an agent are coupled to one antibody molecule. In another embodiment, more than one type of agent may be coupled to one antibody. Regardless of the particular embodiment, immunoconjugates with more than one agent may be prepared in a variety of ways. For example, more than one agent may be coupled directly to an antibody molecule, or linkers which provide multiple sites for attachment can be used. Alternatively, a carrier can be used.

A carrier may bear the agents in a variety of ways, including covalent bonding either directly or via a linker group. Suitable carriers include proteins such as albumins (*e.g.*, U.S. Patent No. 4,507,234, to Kato et al.), peptides and polysaccharides such as aminodextran (*e.g.*, U.S. Patent No. 4,699,784, to Shih et al.). A carrier may also bear an agent by noncovalent bonding or by encapsulation, such as within a liposome vesicle (*e.g.*, U.S. Patent Nos. 4,429,008 and 4,873,088). Carriers specific for radionuclide agents include radiohalogenated small molecules and chelating compounds. For example, U.S. Patent No. 4,735,792 discloses representative radiohalogenated small molecules and their synthesis. A radionuclide chelate may be formed from chelating compounds that include those containing nitrogen and sulfur atoms as the donor atoms for binding the metal, or metal oxide, radionuclide. For example, U.S. Patent No. 4,673,562, to Davison et al. discloses representative chelating compounds and their synthesis.

A variety of routes of administration for the antibodies and immunoconjugates may be used. Typically, administration will be intravenous,

intramuscular, subcutaneous or in the bed of a resected tumor. It will be evident that the precise dose of the antibody/immunoconjugate will vary depending upon the antibody used, the antigen density on the tumor, and the rate of clearance of the antibody.

5 Diagnostic reagents of the present invention may also comprise at least a portion of a polynucleotide disclosed herein. For example, at least two oligonucleotide primers may be employed in a polymerase chain reaction (PCR) based assay to amplify breast tumor-specific cDNA derived from a biological sample, wherein at least one of the oligonucleotide primers is specific for a polynucleotide
10 encoding a breast tumor protein of the present invention. The presence of the amplified cDNA is then detected using techniques well known in the art, such as gel electrophoresis. Similarly, oligonucleotide probes specific for a polynucleotide encoding a breast tumor protein of the present invention may be used in a hybridization assay to detect the presence of an inventive polypeptide in a
15 biological sample.

As used herein, the term "oligonucleotide primer/probe specific for a polynucleotide" means an oligonucleotide sequence that has at least about 60%, preferably at least about 75% and more preferably at least about 90%, identity to the polynucleotide in question, or an oligonucleotide sequence that is anti-sense to a
20 sequence that has at least about 60%, preferably at least about 75% and more preferably at least about 90%, identity to the polynucleotide in question. Oligonucleotide primers and/or probes which may be usefully employed in the inventive diagnostic methods preferably have at least about 10-40 nucleotides. In a preferred embodiment, the oligonucleotide primers comprise at least about 10
25 contiguous nucleotides of a polynucleotide disclosed herein or that is anti-sense to a polynucleotide sequence disclosed herein. Preferably, oligonucleotide probes for use in the inventive diagnostic methods comprise at least about 15 contiguous oligonucleotides of a polynucleotide that encodes one of the polypeptides disclosed herein or that is anti-sense to a sequence that encodes one of the polypeptides
30 disclosed herein. Techniques for both PCR based assays and hybridization assays are

well known in the art (see, for example, Mullis *et al. Ibid*; Ehrlich, *Ibid*). Primers or probes may thus be used to detect breast tumor-specific sequences in biological samples, including blood, urine and/or breast tumor tissue.

The following Examples are offered by way of illustration and not by
5 way of limitation.

EXAMPLES

Example 1

10 ISOLATION AND CHARACTERIZATION OF BREAST
TUMOR POLYPEPTIDES

This Example describes the isolation of breast tumor polypeptides from a breast tumor cDNA library.

15 A human breast tumor cDNA expression library was constructed from a pool of breast tumor poly A⁺ RNA from three patients using a Superscript Plasmid System for cDNA Synthesis and Plasmid Cloning kit (BRL Life Technologies, Gaithersburg, MD 20897) following the manufacturer's protocol. Specifically, breast tumor tissues were homogenized with polytron (Kinematica, Switzerland) and total
20 RNA was extracted using Trizol reagent (BRL Life Technologies) as directed by the manufacturer. The poly A⁺ RNA was then purified using a Qiagen oligotex spin column mRNA purification kit (Qiagen, Santa Clarita, CA 91355) according to the manufacturer's protocol. First-strand cDNA was synthesized using the NotI/Oligo-dT18 primer. Double-stranded cDNA was synthesized, ligated with EcoRI/BstX I
25 adaptors (Invitrogen, Carlsbad, CA) and digested with NotI. Following size fractionation with Chroma Spin-1000 columns (Clontech, Palo Alto, CA 94303), the cDNA was ligated into the EcoRI/NotI site of pCDNA3.1 (Invitrogen, Carlsbad, CA) and transformed into ElectroMax *E. coli* DH10B cells (BRL Life Technologies) by electroporation.

Using the same procedure, a normal human breast cDNA expression library was prepared from a pool of four normal breast tissue specimens. The cDNA libraries were characterized by determining the number of independent colonies, the percentage of clones that carried insert, the average insert size and by sequence analysis. The breast tumor library contained 1.14×10^7 independent colonies, with more than 90% of clones having a visible insert and the average insert size being 936 base pairs. The normal breast cDNA library contained 6×10^6 independent colonies, with 83% of clones having inserts and the average insert size being 1015 base pairs. Sequencing analysis showed both libraries to contain good complex cDNA clones that were synthesized from mRNA, with minimal rRNA and mitochondrial DNA contamination sequencing.

cDNA library subtraction was performed using the above breast tumor and normal breast cDNA libraries, as described by Hara *et al.* (*Blood*, 84:189-199, 1994) with some modifications. Specifically, a breast tumor-specific subtracted cDNA library was generated as follows. Normal breast cDNA library (70 µg) was digested with EcoRI, NotI, and SfuI, followed by a filling-in reaction with DNA polymerase Klenow fragment. After phenol-chloroform extraction and ethanol precipitation, the DNA was dissolved in 100 µl of H₂O, heat-denatured and mixed with 100 µl (100 µg) of Photoprobe biotin (Vector Laboratories, Burlingame, CA), the resulting mixture was irradiated with a 270 W sunlamp on ice for 20 minutes. Additional Photoprobe biotin (50 µl) was added and the biotinylation reaction was repeated. After extraction with butanol five times, the DNA was ethanol-precipitated and dissolved in 23 µl H₂O to form the driver DNA.

To form the tracer DNA, 10 µg breast tumor cDNA library was digested with BamHI and Xhol, phenol chloroform extracted and passed through Chroma spin-400 columns (Clontech). Following ethanol precipitation, the tracer DNA was dissolved in 5 µl H₂O. Tracer DNA was mixed with 15 µl driver DNA and 20 µl of 2 x hybridization buffer (1.5 M NaCl/10 mM EDTA/50 mM HEPES pH 7.5/0.2% sodium dodecyl sulfate), overlaid with mineral oil, and heat-denatured completely. The sample was immediately transferred into a 68 °C water bath and

incubated for 20 hours (long hybridization [LH]). The reaction mixture was then subjected to a streptavidin treatment followed by phenol/chloroform extraction. This process was repeated three more times. Subtracted DNA was precipitated, dissolved in 12 µl H₂O, mixed with 8 µl driver DNA and 20 µl of 2 x hybridization buffer, and 5 subjected to a hybridization at 68 °C for 2 hours (short hybridization [SH]). After removal of biotinylated double-stranded DNA, subtracted cDNA was ligated into BamHI/XhoI site of chloramphenicol resistant pBCSK⁺ (Stratagene, La Jolla, CA 92037) and transformed into ElectroMax *E. coli* DH10B cells by electroporation to generate a breast tumor specific subtracted cDNA library.

10 To analyze the subtracted cDNA library, plasmid DNA was prepared from 100 independent clones, randomly picked from the subtracted breast tumor specific library and characterized by DNA sequencing with a Perkin Elmer/Applied Biosystems Division Automated Sequencer Model 373A (Foster City, CA). Thirty-eight distinct cDNA clones were found in the subtracted breast tumor-specific cDNA library. The determined 3' cDNA sequences for 14 of these clones are provided in SEQ ID NO: 1-14, with the corresponding 5' cDNA sequences being provided in SEQ ID NO: 15-28, respectively. The determined one strand (5' or 3') cDNA sequences for the remaining clones are provided in SEQ ID NO: 29-52. Comparison of these cDNA sequences with known sequences in the gene bank using the EMBL 15 and GenBank databases (Release 97) revealed no significant homologies to the sequences provided in SEQ ID NO: 3, 10, 17, 24 and 45-52. The sequences provided in SEQ ID NO: 1, 2, 4-9, 11-16, 18-23, 25-41, 43 and 44 were found to show at least some degree of homology to known human genes. The sequence of SEQ ID NO: 42 was found to show some homology to a known yeast gene.

20 25 cDNA clones isolated in the breast subtraction described above were colony PCR amplified and their mRNA expression levels in breast tumor, normal breast and various other normal tissues were determined using microarray technology (Synteni, Fremont, CA). Briefly, the PCR amplification products were dotted onto slides in an array format, with each product occupying a unique location in the array. 30 mRNA was extracted from the tissue sample to be tested, reverse transcribed, and

fluorescent-labeled cDNA probes were generated. The microarrays were probed with the labeled cDNA probes, the slides scanned and fluorescence intensity was measured. This intensity correlates with the hybridization intensity.

Data was analyzed using GEMTOOLS Software. Twenty one distinct
5 cDNA clones were found to be over-expressed in breast tumor and expressed at low levels in all normal tissues tested. The determined partial cDNA sequences for these clones are provided in SEQ ID NO: 53-73. Comparison of the sequences of SEQ ID NO: 53, 54 and 68-71 with those in the gene bank as described above, revealed some homology to previously identified human genes. No significant homologies were
10 found to the sequences of SEQ ID NO: 55-67, 72 (referred to as JJ 9434) and 73 (referred to as B535S). In further studies, full length cDNA sequences were obtained for the clones 1016F8 (SEQ ID NO: 56; also referred to as B511S) and 1016D12 (SEQ ID NO: 61; also referred to as B532S), and an extended cDNA sequence was obtained for 1012H8 (SEQ ID NO: 64; also referred to as B533S). These cDNA
15 sequences are provided in SEQ ID NO: 95-97, respectively, with the corresponding predicted amino acid sequences for B511S and B532S being provided in SEQ ID NO: 98 and 99, respectively.

Analysis of the expression of B511S in breast tumor tissues and in a variety of normal tissues (skin, PBMC, intestine, breast, stomach, liver, kidney, fetal
20 tissue, adrenal gland, salivary gland, spinal cord, large intestine, small intestine, bone marrow, brain, heart, colon and pancreas) by microarray, northern analysis and real time PCR, demonstrated that B511S is over-expressed in breast tumors, and normal breast, skin and salivary gland, with expression being low or undetectable in all other tissues tested.

25 Analysis of the expression of B532S in breast tumor tissue and in a variety of normal tissues (breast, PBMC, esophagus, HMEC, spinal cord, bone, thymus, brain, bladder, colon, liver, lung, skin, small intestine, stomach, skeletal muscle, pancreas, aorta, heart, spleen, kidney, salivary gland, bone marrow and adrenal gland) by microarray, Northern analysis and real time PCR, demonstrated that

B532S is over-expressed in 20-30% of breast tumors with expression being low or undetectable in all other tissues tested.

In a further experiment, cDNA fragments were obtained from two subtraction libraries derived by conventional subtraction, as described above and 5 analyzed by DNA microarray. In one instance the tester was derived from primary breast tumors, referred to as Breast Subtraction 2, or BS2. In the second instance, a metastatic breast tumor was employed as the tester, referred to as Breast Subtraction 3, or BS3. Drivers consisted of normal breast.

cDNA fragments from these two libraries were submitted as templates 10 for DNA microarray analysis, as described above. DNA chips were analyzed by hybridizing with fluorescent probes derived from mRNA from both tumor and normal tissues. Analysis of the data was accomplished by creating three groups from the sets of probes, referred to as breast tumor/mets, normal non-breast tissues, and metastatic breast tumors. Two comparisons were performed using the modified Gemtools 15 analysis. The first comparison was to identify templates with elevated expression in breast tumors. The second was to identify templates not recovered in the first comparison that yielded elevated expression in metastatic breast tumors. An arbitrary level of increased expression (mean of tumor expression versus the mean of normal tissue expression) was set at approximately 2.2.

20 In the first round of comparison to identify over-expression in breast tumors, two novel gene sequences were identified, hereinafter referred to as B534S and B538S (SEQ ID NO: 89 and 90, respectively), together with six sequences that showed some degree of homology to previously identified genes (SEQ ID NO: 74-79). The sequences of SEQ ID NO: 75 and 76 were subsequently determined to be 25 portions of B535S (SEQ ID NO: 73). In a second comparison to identify elevated expression in metastatic breast tumors, five novel sequences were identified, hereinafter referred to as B535S, B542S, B543S, P501S and B541S (SEQ ID NO: 73 and 91-94, respectively), as well as nine gene sequences that showed some homology to known genes (SEQ ID NO: 80-88). Clone B534S and B538S (SEQ ID NO: 89 and

90) were shown to be over-expressed in both breast tumors and metastatic breast tumors.

In a subsequent series of studies, 457 clones from Breast Subtraction 2 were analyzed by microarray on Breast Chip 3. As described above, a first comparison to identify over-expression in breast tumors over normal non-breast tissues was performed. This analysis yielded six cDNA clones that demonstrated elevated expression in breast tumor over normal non-breast tissues. Two of these clones, referred to as 1017C2 (SEQ ID NO: 102) and B546S (SEQ ID NO: 107) do not share significant homology to any known genes. Clone B511S also showed over-expression in breast tumor, which was previously described as 1016F8, with the determined cDNA sequence provided in SEQ ID NO: 95 and the predicted amino acid sequence provided in SEQ ID NO: 98. The remaining four clones over-expressed in breast tumor were found to share some degree of homology to Tumor Expression Enhanced Gene (SEQ ID NO: 103 and 104) Stromelysin-3 (SEQ ID NO: 105) or Collagen (SEQ ID NO: 106).

In the second comparison to determine genes with elevated expression in metastatic breast tumors over non-breast normal tissues, a profile similar to the first comparison was derived. The two putatively novel clones, 1017C2 and B546S, SEQ ID NO: 102 and 107, respectively, were overexpressed in metastatic breast tumors. In addition, Tumor Expression Enhanced Gene and B511S also showed elevated expression in metastatic breast tumors.

As described in U.S. Patent Application No. 08/806,099, filed February 25, 1997, the antigen P501S was isolated by subtracting a prostate tumor cDNA library with a normal pancreas cDNA library and with three genes found to be abundant in a previously subtracted prostate tumor specific cDNA library: human glandular kallikrein, prostate specific antigen (PSA), and mitochondria cytochrome C oxidase subunit II. The determined full-length cDNA sequence for P501S is provided in SEQ ID NO: 100, with the corresponding predicted amino acid sequence being provided in SEQ ID NO: 101. Expression of P501S in breast tumor was examined by microarray analysis. Over-expression was found in prostate tumor, breast tumor

and metastatic breast tumor, with negligible to low expression being seen in normal tissues. This data suggests that P501S may be over-expressed in various breast tumors as well as in prostate tumors.

5

Example 2

GENERATION OF HUMAN CD8+ CYTOTOXIC T-CELLS THAT RECOGNIZE ANTIGEN PRESENTING CELLS EXPRESSING BREAST TUMOR ANTIGENS

- 10 This Example illustrates the generation of T cells that recognize target cells expressing the antigen B511S, also known as 1016-F8 (SEQ ID NO: 56). Human CD8+ T cells were primed *in-vitro* to the B511S gene product using dendritic cells infected with a recombinant vaccinia virus engineered to express B511S as follows (also see Yee et al., Journal of Immunology (1996) 157 (9):4079-86).
- 15 Dendritic cells (DC) were generated from peripheral blood derived monocytes by differentiation for 5 days in the presence of 50 µg/ml GMCSF and 30 µg/ml IL-4. DC were harvested, plated in wells of a 24-well plate at a density of 2×10^5 cells/well and infected for 12 hours with B511S expressing vaccinia at a multiplicity of infection of 5. DC were then matured overnight by the addition of 3 µg/ml CD40-
- 20 Ligand and UV irradiated at 100µW for 10 minutes. CD8+ T cells were isolated using magnetic beads, and priming cultures were initiated in individual wells (typically in 24 wells of a 24-well plate) using 7×10^5 CD8+ T cells and 1×10^6 irradiated CD8-depleted PBMC. IL-7 at 10 ng/ml was added to cultures at day 1. Cultures were re-stimulated every 7-10 days using autologous primary fibroblasts
- 25 retrovirally transduced with B511S and the costimulatory molecule B7.1. Cultures were supplemented at day 1 with 15 I.U. of IL-2. Following 4 such stimulation cycles, CD8+ cultures were tested for their ability to specifically recognize autologous fibroblasts transduced with B511S using an interferon- γ Elispot assay (see Lalvani et al J. Experimental Medicine (1997) 186:859-965). Briefly, T cells from
- 30 individual microcultures were added to 96-well Elispot plates that contained autologous fibroblasts transduced to express either B511S or as a negative control

- antigen EGFP, and incubated overnight at 37° C; wells also contained IL-12 at 10 ng/ml. Cultures were identified that specifically produced interferon- γ only in response to B511S transduced fibroblasts; such lines were further expanded and also cloned by limiting dilution on autologous B-LCL retrovirally transduced with B511S.
- 5 Lines and clones were identified that could specifically recognize autologous B-LCL transduced with B511S but not autologous B-LCL transduced with the control antigens EGFP or HLA-A3. An example demonstrating the ability of human CTL cell lines derived from such experiments to specifically recognize and lyse B511S expressing targets is presented in Figure 1.

10

Example 3
SYNTHESIS OF POLYPEPTIDES

- Polypeptides may be synthesized on an Perkin Elmer/Applied Biosystems Division 430A peptide synthesizer using Fmoc chemistry with HPTU (O-Benzotriazole-N,N,N',N'-tetramethyluronium hexafluorophosphate) activation. A Gly-Cys-Gly sequence may be attached to the amino terminus of the peptide to provide a method of conjugation, binding to an immobilized surface, or labeling of the peptide. Cleavage of the peptides from the solid support may be carried out using
- 15 the following cleavage mixture: trifluoroacetic acid:ethanedithiol:thioanisole:water:phenol (40:1:2:2:3). After cleaving for 2 hours, the peptides may be precipitated in cold methyl-t-butyl-ether. The peptide pellets may then be dissolved in water containing 0.1% trifluoroacetic acid (TFA) and lyophilized prior to purification by C18 reverse phase HPLC. A gradient of 0%-60%
- 20 acetonitrile (containing 0.1% TFA) in water (containing 0.1% TFA) may be used to elute the peptides. Following lyophilization of the pure fractions, the peptides may be characterized using electrospray or other types of mass spectrometry and by amino acid analysis.
- 25

From the foregoing, it will be appreciated that, although specific embodiments of the invention have been described herein for the purposes of illustration, various modifications may be made without deviating from the spirit and scope of the invention.

CLAIMS

1. An isolated polypeptide comprising an immunogenic portion of a breast protein or a variant thereof, wherein said protein comprises an amino acid sequence encoded by a polynucleotide comprising a sequence selected from the group consisting of: (a) nucleotide sequences recited in SEQ ID NOS: 3, 10, 17, 24, 45-52, 55-67, 72, 73, 89-97, 102 and 107; (b) complements of said nucleotide sequences; and (c) sequences that hybridize to a sequence of (a) or (b) under moderately stringent conditions.
2. The isolated polypeptide of claim 1, wherein the polypeptide comprises an amino acid sequence selected from the group consisting of SEQ ID NO: 98, 99 and 101.
3. An isolated polynucleotide comprising a nucleotide sequence encoding the polypeptide of any one of claims 1 and 2.
4. An isolated polynucleotide comprising a sequence provided in SEQ ID NOS: 3, 10, 17, 24, 45-52, 55-67, 72, 73, 89-97, 102 and 107.
5. An expression vector comprising a polynucleotide according to any one of claims 3 and 4.
6. A host cell transformed with the expression vector of claim 5.
7. The host cell of claim 6 wherein the host cell is selected from the group consisting of *E. coli*, yeast and mammalian cell lines.
8. A pharmaceutical composition comprising the polypeptide of claim 1 and a physiologically acceptable carrier.
9. A vaccine comprising the polypeptide of claim 1 and a non-specific immune response enhancer.

10. The vaccine of claim 9 wherein the non-specific immune response enhancer is an adjuvant.

11. A vaccine comprising an isolated polynucleotide of any one of claims 3 and 4, and a non-specific immune response enhancer.

12. The vaccine of claim 11 wherein the non-specific immune response enhancer is an adjuvant.

13. A pharmaceutical composition for the treatment of breast cancer comprising a polypeptide and a physiologically acceptable carrier, the polypeptide comprising an immunogenic portion of a breast protein, wherein said protein comprises an amino acid sequence encoded by a polynucleotide comprising a sequence selected from the group consisting of: (a) nucleotide sequences recited in SEQ ID NOS: 1, 2, 4-9, 11-16, 18-23, 25-44, 53, 54, 68-71, 74-88 and 103-106; (b) complements of said nucleotide sequences; and (c) sequences that hybridize to a sequence of (a) or (b) under moderately stringent conditions.

14. A vaccine for the treatment of breast cancer comprising a polypeptide and a non-specific immune response enhancer, said polypeptide comprising an immunogenic portion of a breast protein, wherein said protein comprises an amino acid sequence encoded by a polynucleotide comprising a sequence selected from the group consisting of: (a) nucleotide sequences recited in SEQ ID NOS: 1, 2, 4-9, 11-16, 18-23, 25-44, 53, 54, 68-71, 74-88 and 103-106; (b) complements of said nucleotide sequences; and (c) sequences that hybridize to a sequence of (a) or (b) under moderately stringent conditions.

15. The vaccine of claim 14 wherein the non-specific immune response enhancer is an adjuvant.

16. A vaccine for the treatment of breast cancer comprising a polynucleotide and a non-specific immune response enhancer, the polynucleotide

comprising a sequence selected from the group consisting of: (a) nucleotide sequences recited in SEQ ID NOS: 1, 2, 4-9, 11-16, 18-23, 25-44, 53, 54, 68-71, 74-88 and 103-106; (b) complements of said nucleotide sequences; and (c) sequences that hybridize to a sequence of (a) or (b) under moderately stringent conditions.

17. The vaccine of claim 16, wherein the non-specific immune response enhancer is an adjuvant.

18. A pharmaceutical composition according to any one of claims 8 and 13, for use in the manufacture of a medicament for inhibiting the development of breast cancer in a patient.

19. A vaccine according to any one of claims 9, 11, 14 or 16, for use in the manufacture of a medicament for inhibiting the development of breast cancer in a patient.

20. A fusion protein comprising at least one polypeptide according to claim 1.

21. A pharmaceutical composition comprising a fusion protein according to claim 20 and a physiologically acceptable carrier.

22. A vaccine comprising a fusion protein according to claim 20 and a non-specific immune response enhancer.

23. The vaccine of claim 22 wherein the non-specific immune response enhancer is an adjuvant.

24. A pharmaceutical composition according to claim 21, for use in manufacture of a medicament for inhibiting the development of breast cancer in a patient.

25. A vaccine according to claim 22, for use in the manufacture of a medicament for inhibiting the development of breast cancer in a patient.

26. A method for detecting breast cancer in a patient, comprising:

(a) contacting a biological sample from a patient with a binding agent which is capable of binding to a polypeptide, the polypeptide comprising an immunogenic portion of a breast protein, wherein said protein comprises an amino acid sequence encoded by a polynucleotide comprising a sequence selected from the group consisting of nucleotide sequences recited in SEQ ID NOS: 1-97, 100 and 102-107, complements of said nucleotide sequences and sequences that hybridize to a sequence provided in SEQ ID NO: 1-97, 100 and 102-107 under moderately stringent conditions; and

(b) detecting in the sample a protein or polypeptide that binds to the binding agent, thereby detecting breast cancer in the patient.

27. The method of claim 26 wherein the binding agent is a monoclonal antibody.

28. The method of claim 27 wherein the binding agent is a polyclonal antibody.

29. A method for monitoring the progression of breast cancer in a patient, comprising:

(a) contacting a biological sample from a patient with a binding agent that is capable of binding to a polypeptide, said polypeptide comprising an immunogenic portion of a breast protein, wherein said protein comprises an amino acid sequence encoded by a polynucleotide comprising a sequence selected from the group consisting of nucleotide sequences recited in SEQ ID NOS: 1-97, 100 and 102-107, complements of said nucleotide sequences and sequences that hybridize to a sequence provided in SEQ ID NO: 1-97, 100 and 102-107 under moderately stringent conditions;

(b) determining in the sample an amount of a protein or polypeptide that binds to the binding agent;

- (c) repeating steps (a) and (b); and
- (d) comparing the amount of polypeptide detected in steps (b) and (c)

to monitor the progression of breast cancer in the patient.

30. A monoclonal antibody that binds to a polypeptide comprising an immunogenic portion of a breast protein or a variant of said protein that differs only in conservative substitutions and/or modifications, wherein said protein comprises an amino acid sequence encoded by a polynucleotide comprising a sequence selected from the group consisting of: (a) nucleotide sequences recited in SEQ ID NOS: 3, 10, 17, 24, 45-52, 55-67, 72, 73, 89-97, 102 and 107; (b) complements of said nucleotide sequences; and (c) sequences that hybridize to a sequence of (a) or (b) under moderately stringent conditions.

31. A monoclonal antibody according to claim 30, for use in the manufacture of a medicament for inhibiting the development of breast cancer in a patient.

32. The monoclonal antibody of claim 31 wherein the monoclonal antibody is conjugated to a therapeutic agent.

33. A method for detecting breast cancer in a patient comprising:
(a) contacting a biological sample from a patient with at least two oligonucleotide primers in a polymerase chain reaction, wherein at least one of the oligonucleotides is specific for a polynucleotide encoding a polypeptide comprising an immunogenic portion of a breast protein, said protein comprising an amino acid sequence encoded by a polynucleotide comprising a sequence selected from the group consisting of nucleotide sequences recited in SEQ ID NO: 1-97, 100 and 102-107, complements of said nucleotide sequences and sequences that hybridize to a sequence of SEQ ID NO: 1-97, 100 and 102-107 under moderately stringent conditions; and
(b) detecting in the sample a polynucleotide sequence that amplifies in the presence of the oligonucleotide primers, thereby detecting breast cancer.

34. The method of claim 33, wherein at least one of the oligonucleotide primers comprises at least about 10 contiguous nucleotides of a polynucleotide comprising a sequence selected from SEQ ID NOS: 1-97, 100 and 102-107.

35. A diagnostic kit comprising:

- (a) one or more monoclonal antibodies of claim 30; and
- (b) a detection reagent.

36. A diagnostic kit comprising:

(a) one or more monoclonal antibodies that bind to a polypeptide encoded by a polynucleotide comprising a nucleotide sequence selected from the group consisting of SEQ ID NOS: 1, 2, 4-9, 11-16, 18-23, 25-44, 53, 54, 68-71, 74-88 and 103-106, complements of said sequences, and sequences that hybridize to a sequence of SEQ ID NO: 1, 2, 4-9, 11-16, 18-23, 25-44, 53, 54, 68-71, 74-88 or 103-106 under moderately stringent conditions; and

- (b) a detection reagent.

37. The kit of claims 35 or 36 wherein the monoclonal antibodies are immobilized on a solid support.

38. The kit of claim 37 wherein the solid support comprises nitrocellulose, latex or a plastic material.

39. The kit of claims 35 or 36 wherein the detection reagent comprises a reporter group conjugated to a binding agent.

40. The kit of claim 39 wherein the binding agent is selected from the group consisting of anti-immunoglobulins, Protein G, Protein A and lectins.

41. The kit of claim 39 wherein the reporter group is selected from the group consisting of radioisotopes, fluorescent groups, luminescent groups, enzymes, biotin and dye particles.

42. A diagnostic kit comprising at least two oligonucleotide primers, at least one of the oligonucleotide primers being specific for a polynucleotide encoding a polypeptide comprising an immunogenic portion of a breast protein, said protein comprising an amino acid sequence encoded by a polynucleotide comprising a sequence selected from the group consisting of nucleotide sequences recited in SEQ ID NOS: 1-97, 100 and 102-107 complements of said nucleotide sequences and sequences that hybridize to a sequence of SEQ ID NO: 1-97, 100 and 102-107 under moderately stringent conditions.

43. A diagnostic kit of claim 42 wherein at least one of the oligonucleotide primers comprises at least about 10 contiguous nucleotides of a polynucleotide comprising a sequence selected from SEQ ID NOS: 1-97, 100 and 102-107.

44. A method for detecting breast cancer in a patient, comprising:

- (a) obtaining a biological sample from the patient;
- (b) contacting the biological sample with an oligonucleotide probe specific for a polynucleotide encoding a polypeptide comprising an immunogenic portion of a breast protein, said protein comprising an amino acid sequence encoded by a polynucleotide comprising a sequence selected from the group consisting of nucleotide sequences recited in SEQ ID NOS: 1-97, 100 and 102-107, complements of said nucleotide sequences and sequences that hybridize to a sequence of SEQ ID NO: 1-97, 100 and 102-107 under moderately stringent conditions; and
- (c) detecting in the sample a polynucleotide sequence that hybridizes to the oligonucleotide probe, thereby detecting breast cancer in the patient.

45. The method of claim 44 wherein the oligonucleotide probe comprises at least about 15 contiguous nucleotides of a polynucleotide comprising a sequence selected from the group consisting of SEQ ID NOS: 1-97, 100 and 102-107.

46. A diagnostic kit comprising an oligonucleotide probe specific for a polynucleotide encoding a polypeptide comprising an immunogenic portion of a breast protein, said protein comprising an amino acid sequence encoded by a polynucleotide comprising a sequence selected from the group consisting of nucleotide sequences recited in SEQ ID NOS: 1-97, 100 and 102-107, complements of said nucleotide sequences, and sequences that hybridize to a sequence of SEQ ID NO: 1-97, 100 and 102-107 under moderately stringent conditions.

47. The diagnostic kit of claim 46, wherein the oligonucleotide probe comprises at least about 15 contiguous nucleotides of a polynucleotide comprising a sequence selected from the group consisting of SEQ ID NOS: 1-97, 100 and 102-107.

48. A method for treating breast cancer in a patient, comprising the steps of:

- (a) obtaining peripheral blood cells from the patient;
- (b) incubating the cells in the presence of at least one polypeptide of any one of claims 1 and 2, such that T cells proliferate; and administering the proliferated T cells to the patient.

49. A method for treating breast cancer in a patient, comprising the steps of:

- (a) obtaining peripheral blood cells from the patient;
- (b) incubating the cells in the presence of at least one polynucleotide of any one of claims 3 and 4, such that T cells proliferate; and
- (c) administering the proliferated T cells to the patient.

50. The method of any one of claims 48 and 49 wherein the step of incubating the cells is repeated one or more times.

51. The method of any one of claims 48 and 49 wherein step (a) further comprises separating the T cells from the peripheral blood cells and the cells incubated in step (b) are the T cells.

52. The method of any one of claims 48 and 49 wherein step (a) further comprises separating CD4+ cells or CD8+ cells from the peripheral blood cells and the cells proliferated in step (b) are CD4+ or CD8+ T cells.

53. The method of any one of claims 48 and 49 wherein step (b) further comprises cloning at least one T cell that proliferated in the presence of the polypeptide.

54. A composition for the treatment of breast cancer in a patient, comprising T cells proliferated in the presence of a polypeptide of any one of claims 1 and 2, in combination with a pharmaceutically acceptable carrier.

55. A composition for the treatment of breast cancer in a patient comprising T cells proliferated in the presence of a polynucleotide of any one of claims 3 and 4, in combination with a pharmaceutically acceptable carrier.

56. A method for treating breast cancer in a patient, comprising the steps of:

- (a) incubating antigen presenting cells in the presence of at least one polypeptide of any one of claims 1 and 2; and
- (b) administering to the patient the incubated antigen presenting cells.

57. A method for treating breast cancer in a patient, comprising the steps of:

- (a) incubating antigen presenting cells in the presence of at least one polynucleotide of any one of claims 3 and 4; and
- (b) administering to the patient the incubated antigen presenting

cells.

58. The method of claims 56 or 57 wherein the antigen presenting cells are selected from the group consisting of dendritic cells, macrophage cells, monocyte cells, fibroblast cells, B-cells or combinations thereof.

59. A composition for the treatment of breast cancer in a patient, comprising antigen presenting cells incubated in the presence of a polypeptide of any one of claims 1 and 2, in combination with a pharmaceutically acceptable carrier.

60. A composition for the treatment of breast cancer in a patient, comprising antigen presenting cells incubated in the presence of a polynucleotide of any one of claims 3 and 4, in combination with a pharmaceutically acceptable carrier.

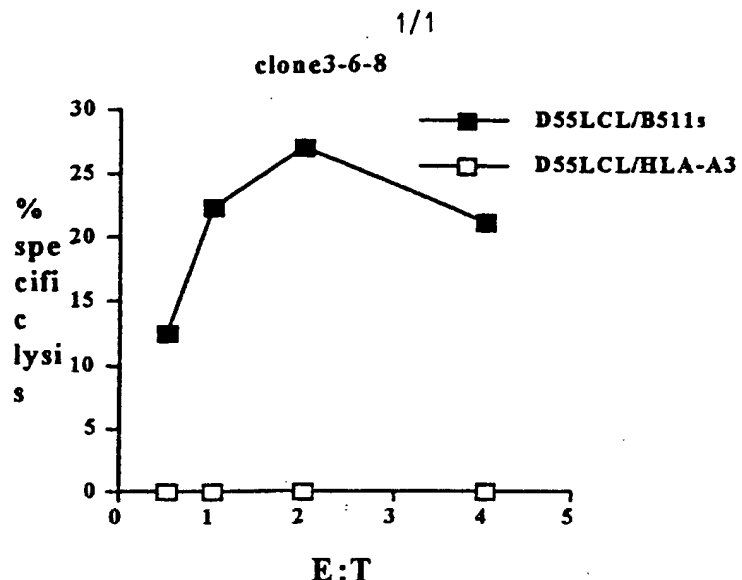
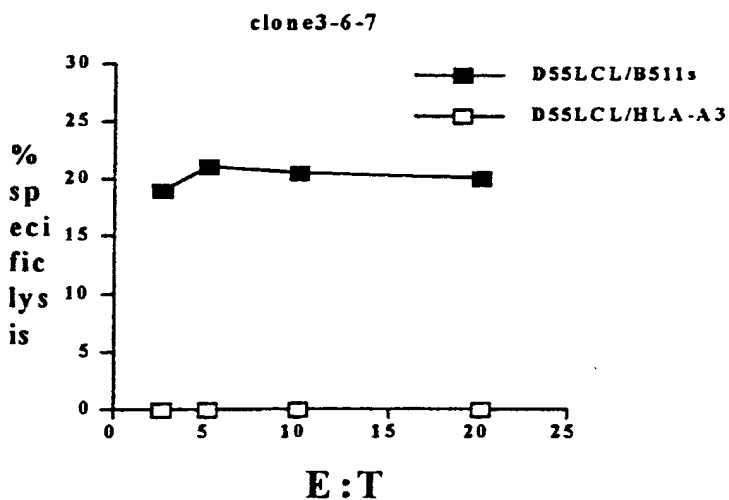
**FIGURE 1A****FIGURE 1B**

Figure 1: Specific lytic activity of B511s-specific CTL clones 3-6-8 and 3-6-7 measured on autologous LCL transduced with B511s (filled squares) or HLA-A3 (open squares). Each data point is the average of triplicate measurements.

FIGS. 1A AND 1B

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Reed, Steven G.
Xu, Jiangchun
Dillon, Davin C.

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DIAGNOSIS OF BREAST CANCER AND METHODS FOR THEIR USE

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tccaggcatc agttggatga ttcatcatgg taattatggc attatcatat tcttcataact
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tggaatattt cactctagac cagaaacagc tcccggtgct ccctcatttt ctgaggctta
aatttn 60
120
180
240
300
360
366

<210> 22
<211> 315
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(315)
<223> n = A,T,C or G

<400> 22
acttaatgca atctctggag gataatttgg atcaagaataa aaagaanaaaa tgaatttagga
gaagaaatna ctgggtnata tttcaatatt ttagaacttt aanaatgtt actatgattt
caatataattt gtaaaaactg agatacangt ttgacctata tctgcatttt gataattaaa
cnaatnnatt ctattnaat gttgttttag agtcacagca cagactgaaa ctttttttga
atacctnaat atcacacttn tncttnaat gatgttgaag acaatgatga catgccttna
gcatataatg tcgac 60
120
180
240
300
315

<210> 23

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<211> 202
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(202)
<223> n = A,T,C or G

<400> 23

actaatccag tgggtgnaa ttccattgtt ttggcaact caggatatta aatttatnat	60
ttaaaaattc ccaagagaaa naaaactccag gccctgattt gttcactggg gaatttacc	120
aatgttnca nnaaganatg acgctgattc tgtnaaatct ttttcagaag atagaggaga	180
acacccaccg ntcatttta tg	202

<210> 24

<211> 365
<212> DNA
<213> Homo sapien

<220>

<221> misc_feature
<222> (1)...(365)
<223> n = A,T,C or G

<400> 24

ggatttcttg ccctttctc ctttttaag tatcaatgtt tgaaatccac ctgtaccacc	60
ctttctgcca tacaaccgct accacatctg gctccttagaa cctgtttgc tttcatagat	120
ggatctcgga accnagtgtt nactcattt taaaacccca ttttagcaga tngttgctn	180
tggctgtct gtattcacca tggggcctgt acacaccacg tgtggttata gtcaaacaca	240
gtgccctcca ttgtggccac atgggagacc catnaccnna tactgcatcc tgggctgatn	300
acggcaactgc atctnacccg acntggatt gaacccgggg tggcagcng aattgaacag	360
gatca	365

<210> 25

<211> 359
<212> DNA
<213> Homo sapien

<220>

<221> misc_feature
<222> (1)...(359)
<223> n = A,T,C or G

<400> 25

gtttctctgt tcaacagtgc ttggacggaa cccggcgctc gttccccacc ccggccggcc	60
gcccatacgcc agccctccgt cacctttca ccgcaccctc ggactgcccc aaggcccccg	120
ccgcctctcc ngcgcncgc agccaccgcc gccnnccnca cctctccctn gtccgcctn	180
nacaacgcgt ccacctcgca ngtcgccng aactaccacc nggactcata ngccgcctc	240
aaccggccga tcaacctgga gctctncccc ccgacntaa cctttccntg tcttacttac	300
nttaaccgccc gnttattttt cttnaaaaaga actttcccc aataactttct ttcaccnnt	359

<210> 26

<211> 400
<212> DNA

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<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(400)
<223> n = A,T,C or G

<400> 26
agtgaaacag tataatgtcaa aaggagtttgcg tgannagcta cataaaaata ttagatatct      60
ttataatttc caataggata ctcatcaattt ttgaataana gacatattct agagaaacca      120
ggtttctgggt ttcagatttg aactctcaag agcttggaaag ttatcactcc catcctcacg      180
acnacnaana aatctnaacn aacnagaanac caatgacttt tcttagatct gtcaaagaac      240
ttcagccacg aggaaaacta tcnccctnaa tactggggac tggaaagaga gggtacagag      300
aatcacagtg aatcatagcc caagatcagc ttgcccggag cttaagctng tacatnatt      360
acttacaggg accacttcac agtnngtnga tnaantgccn      400

<210> 27
<211> 366
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(366)
<223> n = A,T,C or G

<400> 27
gaatttctta gaaactgaag tttactctgt tccaagatata atttcactg tcttaatcaa      60
agggcgctng aatcatagca aatattctca tcttcacta aactttaagt agttntcctg      120
gaattttaca ttttccagaa aacactccct tctgtatctg tgaaagaaag tgtgcctcag      180
gctgttagact gggctgcact ggacacctgc gggggactct ggctnagtgn ggacatggtc      240
agtattgtt ttcctcanac tcagcctgtg tagctntgaa agcatggAAC agattacact      300
gcagtttacg tcatccccaca catcttggac tccnagaccc ggggagggtca catagtccgt      360
tatgna      366

<210> 28
<211> 402
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(402)
<223> n = A,T,C or G

<400> 28
agtgggagcc tcctccttcc ccactcagtt ctttacatcc ccgaggcgca gctgggcnaa      60
ggaagtggcc agctgcagcg cctcctgcag gcagccaaacg ttcttgccctg tggcctgtgc      120
agacacatcc ttgccaccac ctttaccgtc catcangct gacacctgt gcacccactc      180
gctngcttt aagccccgat nggctgcatt ctgggggact tgacacaggc ncgtgatctt      240
gccagccctca ttgtccaccg tgaagagcat ggccaaaaagt ctgagggag tgcatcttga      300
anagttcaa ggcttcattc agggccttng cttaggcgc nctctccatc tccnnggata      360
acnagaggct ggtnnggtn actntcaata aactgcttcg tc      402

<210> 29

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<211> 175
<212> DNA
<213> Homo sapien

<400> 29
cgacgggca tgaccggtcc ggtcagctgg gtggccagtt tcagttcttc agcagaactg      60
tctcccttct tggggggccga gggcttcctg gggaaagagga tgagtttggg gcggtactcc      120
ttcagccgct gcacgttggt ctgcagggac tccgtggact tggccgcct cctcg      175

<210> 30
<211> 360
<212> DNA
<213> Homo sapien

<400> 30
ttgttatttct tatgtatctct gatgggttct tctcgaaaat gccaaagtggg agacttttg      60
gcatgctcca gatttaaatc cagctgaggc tccctttgtt ttcaagttcca tgtaacaatc      120
tggaaaggaaa cttcacggac aggaagactg ctggagaaga gaagcgtgtt agcccatgg      180
aggtctgggg aatcatgtaa agggtaccca gacctcactt ttagtttattt acatcaatga      240
gttcttcag ggaaccaaac ccagaattcg gtgcaaaagc caaacatctt ggtgggattt      300
gataaatgcc ttgggacctg gagtgctggg cttgtgcaca ggaagagcac cagccgctga      360

<210> 31
<211> 380
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(380)
<223> n = A,T,C or G

<400> 31
acgctctaag cctgtccacg agctcaatag ggaaggctgt gatgactaca gactttgcga      60
acgctacgcc atggtttatg gatacaatgc tgcttataan cgctacttca ggaagcgccg      120
agggaccnna tgagactgag ggaagaaaaaa aaatctctt ttttctggag gctggcacct      180
gatttgtat cccccctgtttt cagcatncn gaaatacata ggcttatata caatgtttt      240
ttcctgtata ttctcttgta tggctgcacc cttttccc gccccagat tgataagtaa      300
tgaaagtgc ctgcagtnag ggtcaangga gactcancat atgtgattgt tccntnataa      360
acttctggtg tgataactttc      380

<210> 32
<211> 440
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(440)
<223> n = A,T,C or G

<400> 32
gtgtatggga gcccctgact cctcacgtgc ctgatctgtg cccttggccc caggtcaggc      60
ccacccctg cacctccacc tgccccagcc cctgcctctg ccccaagtgg ggccagctgc      120
cctcacttct ggggtggatg atgtgaccctt cctngggggc ctgcggaaagg gacaagggtt      180

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ccctgaagtc ttacggtcca acatcaggac caagtcccat ggacatgctg acagggtccc	240
caggggagac cgtntcanta gggatgtgtg cctggctgtg tacgtgggtg tgcagtgcac	300
gtganaagca cgtggcggtc tctggggcc atgtttgggg aaggaagtgt gcccnccacc	360
cttggagaac ctcagtcnnn gtagccccct gcccctggcac agcngcatnc acttcaaggg	420
cacccttgg gggttgggggt	440
<210> 33	
<211> 345	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1)...(345)	
<223> n = A,T,C or G	
<400> 33	
tatTTtaaca atgtttatta ttcatttatac cctctataga accaccaccc acaccgagga	60
gattatttgg agtgggtccc aaccttagggc ctggactctg aaatctaact ccccaacttcc	120
ctcattttgt gacttaggtg ggggcatggt tcagtcagaa ctggtgtctc ctattggatc	180
gtgcagaagg agacacctgg cacacacata tggtgccac acccaggagg gttgattggc	240
aggcttggaaag aaaaaagtct cccaaataaaag gcactttac ctcaaagang gggtgggaggt	300
tggtctgctg ggaatgttgt tgTTgggggt gggaaagantt atttc	345
<210> 34	
<211> 440	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1)...(440)	
<223> n = A,T,C or G	
<400> 34	
tgtatTTTT ttattggaaa acaaataatac aacttggaat ggatttttag gcaaattgtg	60
ccataaaggc atTTtaagtg gctaaacaaa gttaaaaag caagtaacaa taaaagaaaa	120
tgtttctgtt acaggaccag cagtacaaaa aaatagtgtt cgagtacctg gataatacac	180
ccgttttgc aatgtgcac ttTTtaagtac atattgttga ctgtccatag tccacgcaga	240
gttacaactc cacacttcaa caacaacatg ctgacagttc ctAAAGAAAA ctactttaaa	300
aaaggcataa cccagatgtt ccctcatttgc accaacttcca tctnagtta gatgtgcaga	360
agggcttana ttTCCCAGA gtaagccnca tgcaacatgt tacttgatca atTTCTAAA	420
ataaggTTT aggacaatga	440
<210> 35	
<211> 540	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1)...(540)	
<223> n = A,T,C or G	
<400> 35	

atagatggaa ttattaaga tttcacatg tgatagcaca tagtttaat tgcaccaaa	60
gtactaaca aaactctagc aatcaagaat ggcagcatgt tatttataa caatcaacac	120
ctgtggctt taaaattgg tttcataag ataattata ctgaagtaaa tctagccatg	180
ctttaaaaa atgctttagg tcactccaag cttggcagg aacatttgc ataaacaata	240
ataaaacaat cacaattaa taaataacaa atacaacatt gttaggcata atcatataca	300
gtataaggga aaaggtggta gtgttganta agcagttt agaatagaat accttggcct	360
ctatgaaat atgtctagac actttgattc actcagccct gacattcagt tttcaaagtt	420
aggaaacagg ttctacagta tcatttaca gttccaaca cattgaaaac aagttagaaaa	480
tgtgantt attttatta atgcattaca tcctcaagan ttatcaccaa cccctcaggt	540
<210> 36	
<211> 555	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1)...(555)	
<223> n = A,T,C or G	
<400> 36	
cttcgtgtgc ttggaaaattg gagcctgccc ctggccccat aagcccttgt tggttactga	60
gaagtgtata tggggcccaa nctactggc ccagaacaca gagacagcag cccantgcaa	120
tgctgtcgag cattgcaaac gccatgtgtg gaactaggag gagaatatt ccatttggc	180
agaaaccaca gcattggttt tttctactt gtgtgtctgg gggatgaac gcacagatct	240
gtttgacttt gtataaaaaa tagggctccc ccacccccc cnnttctgt tnctttattg	300
tagcantgct gtctgcaagg gagccctan cccctggcag acananctgc ttcaagtgc	360
ctttcccttc tgctaaatgg atgttgatgc actggaggc tttancctg cccttgcac	420
gcncctgctg gaggaagana aaactctgct ggcacatgaccc acagtttctt gactggangc	480
cnntcaaccct cttgggtgaa gccttggctt gaccctgaca tntgcttgg cnctgggtng	540
gnctgggctt cttnaa	555
<210> 37	
<211> 280	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1)...(280)	
<223> n = A,T,C or G	
<400> 37	
ccaccgacta taagaactat gcccctgtt attcctgtac ctgcacatc caacttttc	60
acgtggattt tgcttggatc ttggcaagaa accctaatct ccctccagaa acagtggact	120
ctctaaaaaa tatcctgact tctaataaca ttgatntcaa gaaaatgacg gtcacagacc	180
aggtgaactg ccccnagctc tcgttaaccag gttctacagg gaggctgcac ccactccatg	240
ttntttctgc ttgccttcc cctaccacccccccat	280
<210> 38	
<211> 303	
<212> DNA	
<213> Homo sapien	
<220>	

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<221> misc_feature
<222> (1)...(303)
<223> n = A,T,C or G

<400> 38
catcgagctg gtgttcttgcgtaaa atgggggtcc cttactgcat      60
tatcaaggga aaggcaagac tgggacgtct agtccacagg aagacacgtca ccactgtcgc
cttcacacag gtgaactcggttttggct nagctggtn aagctatcag      120
gaccaattac aatgaacngat acgatnagat ccgcncntcac tgggttagca atgtcctggg
tcctaagtct gtggctcgta tcgcccnaagct cgaanaggcn aangctaaag aacttgccac      180
taa                                              240
                                              300
                                              303

<210> 39
<211> 300
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(300)
<223> n = A,T,C or G

<400> 39
gactcagcgg ctgggtgtcac aagcccagca ctccagggtcc caaggcatt      60
atcaaatccc accaagatnt ttggcttttgcgtaaaatc tgggtttggct tccctnaaag
aactcattga tgtaaatnac taaaatgttgcgttttgcgtaaaatc ctttacatg attccccaga      120
cctcanatgg gctaacacgc ttctcttc cagcagtttgcgttttgcgtaaaatc aagttacctt      180
ccagattgtt acatggaaact gaanacaag ggagccttgcgttttgcgtaaaatc atctggagca      240
                                              300

<210> 40
<211> 318
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(318)
<223> n = A,T,C or G

<400> 40
cccaacacaaa tggctgagga caaatcagtt ctctgtgacc agacatgaga aggttgccaa      60
tgggtgttg ggcgaccaag gccttcccg agtcttcgtc ctctatgagc tctcgcccat
gatggtaag ctgacggaga agcacaggc cttcacccac ttccctgacag gtgtgtgcgc      120
catcattggg ggcatgttca cagtggttgcgttttgcgtaaaatc actcatcgat tcgctcatct accactcagc
acgaggccatc cagaaaaaaaaa ttgatctngg gaagacnacg tagtcacctt cggtntttcc      180
tctgtctcct ttcttccttgcgttttgcgtaaaatc                                              240
                                              300
                                              318

<210> 41
<211> 302
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(302)

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<223> n = A,T,C or G

<400> 41
acttagatgg ggtccgttca ggggataacca gcgttcacat ttttcctttt aagaaagggt      60
cttggcctga atgttccccca tccggacaca ggctgcacgt ctctgttagt gtcaaagctg      120
ccatnaccat ctcggtaacc tactcttact ccacaatgtc tatnttcaact gcagggctct      180
ataatnagtc cataatgtaa atgcctggcc caagacntat ggcctgagtt tatccnaggc      240
ccaaacnatt accagacatt cctcttanat tgaaaacgga tntctttccc ttggcaaaga      300
tc                                         302

<210> 42
<211> 299
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(299)
<223> n = A,T,C or G

<400> 42
cttaataagt ttaaggccaa ggcccggtcc attcttctag caactgacgt tgccagccg      60
ggtttggaca tacctcatgt aaatgtggtt gtoactttg acattccctac ccattccaag      120
gattacatcc atcgagtagg tcgaacagct agagctgggc gctccggaaa ggctattact      180
tttgcacac agtatgtatgtt ggaactcttc cagcgcatacg aacacttnat tgggaagaaa      240
ctaccaggtt ttccaacaca ggtatgatgag gttatgatgc tnacggaacg cgtcgctna      299

<210> 43
<211> 305
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(305)
<223> n = A,T,C or G

<400> 43
ccaacaatgt caagacagcc gtctgtgaca tcccacctcg tggcctcaan atggcagtca      60
ccttcattgg caatagcaca gccntccggg agctcttcaa ggcacatctcg gagcagttca      120
ctgccatgtt ccgcccggaaag gccttctcc actggtagac acggcgagggc atggacaaga      180
tggagttcac cgaggctgag agcaacatga acgacacctgt ctctnagtagt cagcagtacc      240
gggatgccac cgccagaaana ggaggaggat ttccgttnagg aggccgaaga aggaggcctg      300
aggca                                         305

<210> 44
<211> 399
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(399)
<223> n = A,T,C or G

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<400> 44

tttctgtggg ggaaacctga tctcgacnaa attagagaat tttgtcagcg gtatttcggc	60
tggAACAGAA CGAAAACNGA TNAATCTCTG TTTCCTGTAT TAAAGCAACT CGATNCCAG	120
cAGACACAGC TCCNAATTGA TTCCCTCTT NGATTAGCAC AACAGGGAGA AAGAANATGC	180
TTAACGTATT AAGAGCCNGA GACTAAACAG AGCTTGACA TGTATGCTTA GGAAAGAGAA	240
AGAAGCAGCN GCCCGCGNAA TTNGAAGCNG TTCTGTG CNTGGANAAA GAATTGAGC	300
TTCTTTATTA GGCCAACGAA AAACCCCGAA ANANAGGCNT TACNATACT TNGAAAANTC	360
TCCNGCCNNA AAAAGAAAAGA AGCTTNCNGA TTCTTAACC	399

<210> 45

<211> 440

<212> DNA

<213> Homo sapien

<220>

<221> misc_feature

<222> (1)...(440)

<223> n = A,T,C or G

<400> 45

gcggggaggcag aagctaaggc caaaggccaa gagagtggca gtgccagcac tggtgccagt	60
accaggatcca ataacagtgc cagtgcagg gccagcacca gtggtggtt cagtgctgg	120
gccaggcctga ccgcactct cacatttggg ctttcgctg gccttgggg agctgggtgc	180
agcaccaggc gcagctctgg tgcctgtgg ttccctaca agtgagatTT taggtatctg	240
ccttgggttc agtggggaca tctggggctt angggcngg gataaggagc tggatgattc	300
taggaaggcc cangttggag aangatgtgn anagtgtgcc aagacactgc ttttggcatt	360
ttattccctt ctgtttgctg gangtcaatt gacccttnna ntttcttta cttgtgtttt	420
canatatngt taatcctgccc	440

<210> 46

<211> 472

<212> DNA

<213> Homo sapien

<220>

<221> misc_feature

<222> (1)...(472)

<223> n = A,T,C or G

<400> 46

gctctgtaat ttcacatTTT aaaccttccc ttgacctcac attcctttc ggccacctct	60
gtttctctgt tccttctcac agcaaaaact gttcaaaaga gttgttgatt actttcattt	120
ccactttctc accccccatTC tcccctcaat taactctct tcattcccat gatgccatta	180
tgtggctntt attanagtca ccaaccttat tctccaaaac anaagcaaca aggacttga	240
cttctcagca gcaactcagct ctggtncttT aaacaccccc gttacttgct attcctccta	300
cctcataaca atctccttcc cagcctctac tgctgccttc tctgagtct tcccagggtc	360
ctaggctcag atgttagtgc gctcaaccct gctacacaaa gnaatctct gaaaggctgt	420
aaaaatgtcc atncntgtcc tgtgagtgtat ctncangna naataacaaa tt	472

<210> 47

<211> 550

<212> DNA

<213> Homo sapien

<220>

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<221> misc_feature
<222> (1)...(550)
<223> n = A,T,C or G

<400> 47
ccttcctccg cctggccatc cccagcatgc tcatgctgtg catggagtgg tgggcctatg      60
aggtcgggag ctccctcagt ggtctgtatg aggatggatg acggggactg gtggaaacct      120
ggggccctg tctgggtgca aggcgacagc tgttttctt caccaggcat cctccggatg      180
gtggagctgg gcgcgtcagtc catcggttat gaactggcca tcattgtta catggcccct      240
gcagggttca gtgtggctgc cagtgtccgg gtangaaaacg ctctgggtgc tggagacatg      300
gaagcaggca cggaaagtccct ctaccgttcc cctgctgatt acagtgcct ttgctgtanc      360
cttcagtgtc ctgctgttaa gctgttaagga tcacntgggg tacatttta ctaccgaccg      420
agaacatcat taatctggtg gctcagggtgg ttccaattta tgctgtttcc caccttttgc      480
aagcttgc tgctcaggta cacgccaatt ttgaaaagta aacaacgtgc ctggagatgg      540
gaattctgtc

<210> 48
<211> 214
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(214)
<223> n = A,T,C or G

<400> 48
agaaggacat aaacaagctg aacctgccca agacgtgtga tatcagcttc tcagatccag      60
acaacctccct caacttcaag ctggtcatct gtccatna gggcttctac nagagtggga      120
agtttgtt cagtttaag gtggccagg gttacccgca tgatcccccc aaggtgaagt      180
gtgagacnat ggtctatcac cccnacattt acct      214

<210> 49
<211> 267
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(267)
<223> n = A,T,C or G

<400> 49
atctgcctaa aattttattca aataatgaaa atnaatctgt tttaagaaaat tcagtcttt      60
agtttttagg acaaactatgc acaaatagtac gatggagaat tctttttggaa tnaactctag      120
gtngaggaac ttaatccaaac cggagcttt gtgaaggtca gaanacagga gaggaatct      180
tggcaaggaa tgagacnnga gtttgcataat tgcagctaga gtnaatngtt nttaatggga      240
ctgctttgt gtctcccang gaaagtt      267

<210> 50
<211> 300
<212> DNA
<213> Homo sapien

<220>

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<221> misc_feature
<222> (1)...(300)
<223> n = A,T,C or G

<400> 50
gactgggtca aagctgcatg aaaccaggcc ctggcagcaa cctggaaatg gctggagggtg      60
ggagagaacc tgacttctct ttccctctcc ctccctccaac attactggaa ctctgtcctg      120
ttgggatctt ctgagcttgtt ttccctgctg ggtgggacag aggacaaaagg agaaggggagg      180
gtctagaaga ggcagccctt ctttgtcctc tggggtnaat gagcttgacc tanagtagat      240
ggagagacca anagcctctg attttaatt tccataanat gttcnaagta tatntntacc      300

<210> 51
<211> 300
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(300)
<223> n = A,T,C or G

<400> 51
gggtaaaaatc ctgcagcacc cactctggaa aataactgctc ttaattttcc tgaaggtggc      60
cccctatttc tagttggtcc aggattaggg atgtggggta tagggcattt aaatcccttc      120
aagcgctctc caagcaccccc cggcctgggg gtnagttct catcccgcta ctgctgctgg      180
gatcaggtt aataaatggg actcttctgt tctggcctcc aaagcagcct aaaaactgag      240
gggctctgtt agaggggacc tccaccctnn ggaagtccga ggggctnggg aagggtttct      300

<210> 52
<211> 267
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(267)
<223> n = A,T,C or G

<400> 52
aaaatcaact tcntgcatta atanacanat tctanancag gaagtgaana taattttctg      60
cacctatcaa ggaacnnact tgattgcctc tattnaacan atatatcgag ttnctatact      120
tacctgaata ccnccgcata actctcaacc nanatncntc nccatgacac tcnttcttna      180
atgctantcc cgaattcttc attatatcng tcatgttcgn cctgntnata tatcagcaag      240
gtatgtncn taactgcccga nncaang                                267

<210> 53
<211> 401
<212> DNA
<213> Homo sapien

<400> 53
agscttttagc atcatgtaga agcaaactgc acctatggct gagatagggtg caatgaccta      60
caagattttg tgggtttcttag ctgtccagga aaagccatct tcagtcttgc tgacagtcaa      120
agagcaagtg aaaccatttc cagcctaaac tacataaaaag cagccgaacc aatgattaaa      180
gacctctaag gtcctccataat catcattaaa tatgccccaaa ctcattgtga cttttttattt      240

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tatatacagg attaaaatca acattaaatc atcttattta catggccatc ggtgctgaaa	300
ttgagcattt taaatagtac agtaggctgg tatacattag gaaatggact gcactggagg	360
caaatagaaa actaaagaaa ttagatagc tggaaatgct t	401
<210> 54	
<211> 401	
<212> DNA	
<213> Homo sapien	
<400> 54	
ccccacacaa tggataaaaa cacttatagt aaatggggac attcactata atgatctaag	60
aagctacaga ttgtcatagt tgggtttctg ctttacaaaa ttgctccaga tctggaatgc	120
cagtttgacc ttgtcttct ataatatttc cttttttcc cctctttgaa tctctgtata	180
tttgattctt aactaaaatt gttctcttaa atattctgaa tcctggtaat taaaagttt	240
ggtgtatccc ctttacctcc aaggaaagaa ctactagcta caaaaaatata tttggaataa	300
gcattgtttt ggtataaggc acatattttt gttgaagaca ccagactgaa gtaaacagct	360
gtgcaccca ttatttatag ttttgaagt aacaatatgt a	401
<210> 55	
<211> 933	
<212> DNA	
<213> Homo sapien	
<400> 55	
tttactgctt ggcaaaggatc cctgaggcatc agcagagatg ccgagatgaa atcaggAAC	60
tccttagggga tgggtcttctt attacctggg aacaccttag ccagatgcct tacaccacga	120
tgtgcatcaa ggaatgcctc cgccctctacg caccggtagt aaactatccc ggttactcg	180
caaaccatc acctttccag atggacgctc cttacctgca ggaataactg tgTTTatcaa	240
tatTTggct cttcaccacca accccttattt ctgggaagac cctcaggctt ttaacccctt	300
gagattctcc agggaaaattt ctgaaaaat acatccctat gccttcatac cattctcagc	360
tggattaagg aactgcattt ggcagcattt tgccataatt gagtgtaaag tggcagtggc	420
attaactctg cttcgcttca agctggctcc agaccactca aggccacca gctgtcgtca	480
agttgcctca agtccaagaa tggaaatccat gtgtttgaa aaaaagttt ctaattttaa	540
gtcctttcg tataagaattt aakgagacaa tttcctacc aaaggaagaa caaaaggata	600
aatataatac aaaatataatg tatatgttg tttgacaaat tataataactt aggataactt	660
tgactggttt tgacatccat taacagtaat tttaatttct ttgctgtatc tggtaaacc	720
cacaAAAACA cctgaaaaaa ctcaagctga gttccaatgc gaagggaaat gattggttt	780
ggtaactagt ggttagagtgg ctttcaagca tagtttgc taaaactccac tcagtatctg	840
cattactttt atctctgcaat atatctgcat gatacttta ttctcagttt tctttccccca	900
taataaaaaaa tatctgccaat aaaaaaaaaaaa aaa	933
<210> 56	
<211> 480	
<212> DNA	
<213> Homo sapien	
<400> 56	
ggctttaag cattttgtc tttttttttt gatcttcagg tcaccaccat gaagttctta	60
gcagttctgg tacttttggg agtttccatc tttttttttt ctgcccagaa tccgacaaca	120
gctgctccag ctgacacacgtt tccagctact ggtctgtct atgatgaagc ccctgtatct	180
gaaaccactg ctgctgcaac cactgcgacc actgtctgtc ctaccactgc aaccaccgct	240
gcttctacca ctgctcgtaa agacatcca gtttaccca aatgggttgg ggatctcccg	300
aatggtagag tttttttttt agatggaaatc agcttgatgc ttctgcaattt ggtcacaact	360
attcatgctt cttttttttt catccaacta cttaccttgc ctacgatatc ccctttatct	420
ctaattcattt tttttttttt caaaaaaaa ataaactatgtt gcaacaaaaaaa aaaaaaaaaaa	480

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<210> 57
<211> 798
<212> DNA
<213> Homo sapien

<400> 57
agcctacctg gaaagccaac cagtcctcat aatggacaag atccaccagg tcctccgtg 60
gactaacttt gtatatatggg aagtaaaaat agtaacacc ttgcacgacc aaacgaacga 120
agatgaccag agtactcta accccttaga actgttttc ctttgcatac tgcaaatatgg 180
gatgttattt tttcatgag ctctagaaa ttgcacttgc aagtttattt ttgcctcctg 240
tgttactgcc attccttattt acagtatatt tgagtgaatg attatatttt taaaaagtt 300
catggggctt tttgggtgt cctaaactta caaacattcc actcattctg tttgtactg 360
tgattataat tttgtgata atttctggcc tgattgaagg aaatttgaga ggtctgcatt 420
tatatatttt aaatagattt gatagttt taaattgcct ttttcataa ggtatttata 480
aagttatttt ggggtgtctg ggattgtgtg aaagaaaatt agaacccccgc tgtatttaca 540
tttaccttgg tagtttattt gtggatggca gtttctgtt gtttgggga ctgtggtagc 600
tcttggattt ttttgcataat tacagctgaa atctgtgtca tggattaaac tggcttatgt 660
ggctagaata ggaagagaga aaaaatgaaa tgggtgtta ctaattttt actccccatta 720
aaaatttttta atgttaagaa aaccttaaat aaacatgatt gatcaatatg gaaaaaaaaaa 780
aaaaaaaaaaa aaaaaaaaaa 798

<210> 58
<211> 280
<212> DNA
<213> Homo sapien

<400> 58
ggggcagctc ctgaccctcc acagccaccc ggtcagccac cagctggggc aacgagggtg 60
gaggtcccac tgagcctctc gcctgcccccc gccactcgtc tggctgtgt tgatccaagt 120
ccccctgcctg gtccccccaca aggactccca tccaggcccc ctctgcctc ccccttgtca 180
tggaccatgg tcgtgaggaa gggctcatgc cccttattta tgggaaccat ttcattctaa 240
cagaataaac cgagaaggaa accagaaaaaa aaaaaaaaaaa 280

<210> 59
<211> 382
<212> DNA
<213> Homo sapien

<400> 59
aggcggggc agaagctaaa gccaaagccc aagagagtgg cagtgccagc actggtgcca 60
gtaccagtac caataacagt gccagtgcca gtgccagcac cagtggtggc ttcagtgtc 120
gtgccagcct gaccgcccact ctcacatttgc ggctttcgc tggcttggt ggagctgggt 180
ccagcaccag tggcagctct ggtgcctgtg gtttcctcta caagttagat ttttagatatt 240
gttaatctcg ccagtcttcc tcttcaagcc aggtgcattc ctcagaaacc tactcaacac 300
agcactctag gcagccacta tcaatcaattt gaagttgaca ctctgcattt aatctatttgc 360
ccattaaaaaa aaaaaaaaaaaa aa 382

<210> 60
<211> 602
<212> DNA
<213> Homo sapien

<400> 60
tgaagagccg cgccgggtggag ctgctgccccg atggactgc caaccttgcc aagctgcagc 60

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ttgtgggtgga	aatagtgc	cagcgggtca	tccacttggc	gggtcagtgg	gagaagcacc	120
gggtccatc	ctcgtagta	ccgccactcc	gaaagctgca	ggattgcaga	gagctggaat	180
cttctcgacg	gctggcagag	atccaagaac	tgaccagag	tgtccggcgc	gctgctgaag	240
aggcccgacg	gaaggaggag	gtctataagc	agctgatgtc	agagctggag	actctgccc	300
gagatgtgc	ccggctggcc	tacaccagc	gcatcctgga	gatcgtggc	aacatccgga	360
agcagaagga	agagatcacc	aagatcttgt	ctgatacgaa	ggagcttcag	aaggaaatca	420
actccctatc	tgggaagctg	gaccggacgt	ttgcggtgac	tgatgagctt	gtttcaagg	480
atgccaagaa	ggacgatgct	gttcggagg	cctataagta	tctagctgct	ctgcacgaga	540
actgcagcca	gctcatccag	accatcgagg	acacaggcac	catcatcg	gaggttcgag	600
ac						602

<210> 61
<211> 1368
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(1368)
<223> n = A,T,C or G

<400> 61

ccagtggcg	cgcgtataac	gactcaat	agggcgaatt	gggtaccggg	ccccccctcg	60	
agcggccgccc	ctttttttt	tttttttatt	gatcagaatt	caggcttat	tattgagcaa	120	
tgaaaacagc	taaaacttaa	ttccaagcat	gtgtagttaa	agtttgc	aaaatggatatt	180	
gttcacaaaa	cacattcaat	gtttaaacac	tatttat	aagaaca	aaaaatattaa	240	
attgtttgt	tctaaaaaagc	ccat	ttccct	ccaagtctaa	actttgtat	ttgatattaa	300
gcaatgaagt	tat	tttgta	aatctagtt	aacaagcaga	atagcactag	gcagaataaa	360
aaattgcaca	gacgtatgca	at	tttccaag	atagcattct	ttaaattcag	tttcagctt	420
ccaaagattt	gttgc	ccata	atagactaa	acatataatg	atggctaa	aaaataagta	480
tacgaaaat	taaaaaaagga	aatgt	taagtc	cactctaat	ctcataaaag	gtgagagtaa	540
ggatgctaa	gcaaaataaa	tgt	taggttct	tttttctgt	ttccgtt	ttat	600
gcttc	tttga	tat	gcctt	ggttacccat	ttaagtt	ggtt	660
aatgaaaatt	gatcaat	at	acac	ttgtc	ttt	aaattgc	720
ccaaaaaa	ag	gtt	aggc	at	tc	ggaaactt	780
kgktctgata	tggc	agacar	gata	caagtc	ccacc	aggag	840
ggt	atgggc	tg	acaaggta	ttat	gccc	catgg	900
gttttggat	tat	atag	ac	ctag	atc	tttgc	960
acgctaaact	tcca	agg	aa	tct	ttc	tttgc	1020
agttgcattt	ctact	gtt	gt	gt	tttgc	tttgc	1080
agagagaaag	at	ttt	at	gt	tttgc	tttgc	1140
tat	tttgc	at	tttgc	tttgc	tttgc	tttgc	1200
ttggctgatgc	tat	cat	at	tttgc	tttgc	tttgc	1260
acactggact	agt	ggat	cca	ctag	tttgc	tttgc	1320
ttttgttccc	ttt	at	tttgc	tttgc	tttgc	tttgc	1368

<210> 62
<211> 924
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(924)
<223> n = A,T,C or G

<400> 62

caaagggnaca ggaacagctt gnaaaagtact gnacatncctn cctgcaggga ccagccctt	60
gcctccaaaa gcaataggaa atttaaaaaga tttncaactga gaaggggncc acgttnart	120
tntnaatgtn tcargnanar tncctncaa atgncrnctn cactnaactnr gnattgggt	180
tnccgnrtnc mgnactatnt caggttgaa aaactggatc tgccacttat cagttatgtg	240
accttaaaga actccgttaa ttctcagag cctcagttc cttgtctata agttggag	300
aatattaata ctatcatttt tccaaggatt gatgtgaaca ttaatgaggt gaaatgacag	360
atgtgtatca tggttcctaa taaacatcca aaatatagta cttactattt tcattattat	420
tacttgttg aagctaaaga cctcacaata gaatcccatac cagccccacca gacagagyc	480
tgagtttct agtttggaaag agctattaaa taacaacktc tagtgcataat tctatacttg	540
ttatggtcaa gtaactgggc tcagcattt acattcattt gtcctttaag ttcttagcaat	600
gtgaagcagg aactatgatt atattgacta cataaatgaa gaaattgagg ctcagataca	660
ttaagtaatt ctcccagggt cacacagcta gaactggcaa agccctggat tgatccatga	720
tcttccagca ttgaagaatc ataaatgtaa ataactgcaa ggcctttcc tcagaagagc	780
tcctggtgct tgaccaacc cactagact tgcctctac aggggaacat ctgtgggcct	840
gggaatcact gcacgtcgca agagatgtt cttctgatga attattgtt cgtcagtgg	900
tgtgaaggca aaaaaaaaaaaa aaaa	924

<210> 63

<211> 1079
<212> DNA
<213> Homo sapien

<400> 63

agtcccaaga actcaataat ctcttatgtt ttctttgaa gacttatttt aaatattaac	60
tattpcggt cctgaatgga aaaatataaa cattagctca gagacaatgg ggtacctgtt	120
tggaatccag ctggcagcta taagcaccgt tgaaaactct gacaggctt gtgcctttt	180
tattaaatgg cctcacatcc tgaatgcagg aatgtgttcg tttaaataaaa cattaatctt	240
taatgtgaa ttctgaaaac acaaccataa atcatagttt gttttctgt gacaatgatc	300
tagtacatta ttccctccac agcaaacccta ccttccaga aggtggaaat tgtatttgca	360
acaatcaggg caaaaccac acttgaaaag cattttacaa tattatatct aagttgcaca	420
gaagacccca gtgatcacta gggaaatctac cacagtccag ttttcttaat ccaagaaggt	480
caaacttcgg ggaataatgt gtccctcttc tgctgctgct ctgaaaata ttcgatcaaa	540
acgaagttt caagcagcag ttattccaag attagagttt atttgtgtat cccatgtata	600
ctggcaatgt ttagggttgc ccaaaaactc ccagacatcc acaatgtgt tggtaaacc	660
accacatctg gtaacctctc gatcccttag atttgtatct cctgcaaaaata taactgttagc	720
tgactctgga gccttctgca tttctttaa aaccattttt aactgatca ttgcgtccgc	780
agcatgccct ctggtgctct ccaaattggga tgcataagg caaagctcat ttccctgacac	840
attcacatgc acacataaaa gtttctcat cattttggta cttggaaaag gaataatctc	900
ttggctttt aatttcaactc ttgatttctt caacattata gctgtgaaat atccttcttc	960
atgacctgtta ataatctcat aattacttga tctttctt aggtagctat aatatggggg	1020
aataacttcc tgttagaaata tcacatctgg gctgtacaaa gctaagttagg aacacacccc	1079

<210> 64

<211> 1001
<212> DNA
<213> Homo sapien

<400> 64

gaatgtgcaa cgatcaagtc agggtatctg tggtatccac cactttgagc atttatcgat	60
tctatatgtc agaaacattt caagttatct gttctagcaa ggaaatataa aatacttata	120
gttaactatg gcttatctac agtgcaacta aaaacttagat ttatttcctt tccacctgtg	180
ggtttgtatt catttaccac cctttttca ttccctttctt caccacaca ctgtggccgg	240
cctcaggcat atactattct actgtctgtc tctgtaaaggta ttatcatttt agttccacca	300

tatgagagaa	tgcataaaa	gtttttcttt	ccatgtctgg	cttatttcac	ttaacataat	360
gaccccgct	tccatccatg	ttatTTtat	tacccaatag	tgttcataaa	tatataataca	420
cacatataata	ccacattgca	tttgc当地	tattcattga	cggaaactgg	ttaatgttat	480
atcggttgc当地	tttgtggatag	tgctgcaata	aacacgcaag	tggggatata	atttgaagag	540
ttttttgtt	gatgttcctc	caaattttaa	gatttttg	tctatgtttg	tgaaaatggc	600
gttagtattt	tcatagagat	tgcattgaat	ctgttagattg	cttgggttaa	gtatggttat	660
tttgc当地	ttaatTTTTT	cattccatga	agatgagatg	tcttccatt	gtttgtgtcc	720
tctacatTTT	cttcatcaa	agttttgtg	tatTTTgaa	gtagatgtat	ttcaccttat	780
agatcaagt	tattccctaa	atattttatt	tttgc当地	ttgttagatga	aattgccttc	840
ttgatttttt	tttcaactaa	ttcattatta	gtgtatggaa	atgttatgga	tttttatttg	900
ttgggtttta	atcaaaaaact	gtattaaact	tagatTTTT	tgtggagtt	ttaagtttt	960
ctagatataaa	gatcatgaca	tctacaaaaa	aaaaaaaaaa	a		1001

<210> 65

<211> 575

<212> DNA

<213> Homo sapien

<400> 65

acttgatata	aaaaggatat	ccataatgaa	tattttatac	tgcatcctt	acatttagcca	60
ctaaatacgt	tattgcttga	tgaagacctt	tcacagaatc	ctatggattt	cagcatttca	120
cttggctact	tcataccccat	gccttaaaga	ggggcagttt	ctcaaaaagca	gaaacatgcc	180
gccagttctc	aagtttctt	cctaacttca	tttgaatgtt	agggcagctt	gcccccaatg	240
tggggaggtc	cgaacatttt	ctgaattccc	attttcttgt	tgcggctaa	atgacagttt	300
ctgtcattac	ttagattccc	gatctttccc	aaaggtgtt	atttacaag	aggccagcta	360
atagccagaa	atcatgaccc	tgaaagagag	atgaaatttc	aagctgtt	ccagggcagga	420
gctccagtt	ggcaaagggtt	cttgagaatc	agccatttgg	tacaaaaaaag	atttttaaag	480
cttttatgtt	ataccatgga	gccatagaaaa	ggctatggat	tgtttaaagaa	ctattttaaa	540
gtgttccaga	cccaaaaagg	aaaaaaaaaa	aaaaaa			575

<210> 66

<211> 831

<212> DNA

<213> Homo sapien

<400> 66

atggggctcc ttctgtctaa	cagcccacatt gaaatgggtt	aaaagcaagt cagatcagg	60
gatttgtaaa attgtat	tctgtacatg tatgggc	taattcccac caagaaa	120
agaaaattatc ttttagtta	aaacccaatt tcactttc	aaatatcttc caacttatt	180
atgggtgtc actcaattgc	ctatataat atatata	gtgtgtgtgt gtgtgtgc	240
gtgagcgcac gtgtgttat	gcgtgcgc	gtgtgttatc agacatagg	300
ttctaactt tagatagaag	aggagcaaca tctatg	ccaa atactgtca ttctacaat	360
gtgctaattc cagacctaa	tgatactcc	tttaatttaaaaagagtt taaataatt	420
tctatgtgcc tgtaattcc	ttttgagtgc	tgccacaat gtaacatataat tagtgtaaaa	480
gcagatgaaa caaccacgt	ttctaaagtc	tagggattgt gctataatcc ctatttagtt	540
caaaattaac cagaattctt	ccatgtgaaa	tggacccaaac tcataattatt ttatgtaaa	600
tacagagttt taatgcgt	tgacatccca	cagggaaaaa gaatgtctgt agtgggtgac	660
tgttatcaaa tattttatag	aatacaatga	acggtaaca gactggtaac ttgtttgagt	720
tcccatgaca gattgagac	ttgtcaatag	caaatcattt ttgtat	780
tqattnaaaa aacatcatta	aatatctta	aaqtaaaaa aaaaaaaaaa a	831

<210> 67

<211> 590

<212> DNA

<213> Homo sapien

<400> 67
 gtgctctgtg tattttta ctgcattaga cattgaatag taatttgcgt taagatacgc 60
 ttaaaaggctc ttgtgacca tgccccctt tgttagcaata aaatgtttt tacgaaaact 120
 ttctccctgg attagcagtt taaatgaaac agagtcac aatgaaatga gtatttaaaa 180
 taaaaatttg ccttaatgta tcagttcagc tcacaagtat ttaagatga ttgagaagac 240
 ttgaattaaa gaaaaaaaaa ttctcaatca tattttaaa atataagact aaaattgttt 300
 ttaaaacaca ttccaaatag aagtgagtt gaactgacct tatttatact ctttttaagt 360
 ttgttccctt tccctgtgcc tgtgtcaaat cttcaagtct tgctgaaaat acatttgata 420
 caaagtttc ttagttgtg tttagtttt tgcattgtct gttttggct gaagaaccaa 480
 gaaggcagact ttcttttaa aagaattatt tctcttcaa atatttctat ctttttaaaa 540
 aaattccccc ttatggctta tatacctaca tatttaaaaa aaaaaaaaaa 590

<210> 68
 <211> 291
 <212> DNA
 <213> Homo sapien

<220>
 <221> misc_feature
 <222> (1)...(291)
 <223> n = A,T,C or G

<400> 68
 gttccctttt ccggtcggcg tggcttgcg agtggagtgt ccgctgtgcc cgggcctgca 60
 ccatgagcgt cccggccttc atcgacatca gtgaagaaga tcaggctgtc gagttcgtg 120
 cttatctgaa atctaaagga gctgagattt cagaagagaa ctcggaaagg ggacttcatg 180
 ttgattttgc tcaaattttt gaaggctgtg atgtgtgtct gaaggaggat gataaagatg 240
 ttgaaagtgt gatgaacagt ggggnatcct actcttgatc cggaancncna c 291

<210> 69
 <211> 301
 <212> DNA
 <213> Homo sapien

<220>
 <221> misc_feature
 <222> (1)...(301)
 <223> n = A,T,C or G

<400> 69
 tctatgagca tgccaaggct ctgtggagg atgaaggagt gcgtgcctgc tacgaacgct 60
 ccaacgagta ccagctgatt gactgtgccc agtacttctt ggacaagatc gacgtgatca 120
 agcaggctga ctatgtgccg agcgatcagg acctgctcg ctgcccgtgtc ctgacttctg 180
 gaatcttga gaccaagtcc caggtggacn aagtcaactt ccacatgntt gacgtgggtg 240
 gccagcgcga tgaacgcccgc aagtggatcc agtgcttcaa cgatgtgact gccatcatct 300
 t 301

<210> 70
 <211> 201
 <212> DNA
 <213> Homo sapien

<400> 70
 gcggctcttc ctcgggcagc ggaagcggcg cggcggtcg agaagtggcc taaaacttcg 60

gcgttgggtg aaagaaaaatg gcccgaacca agcagactgc tcgtaagtcc accgggtggga	120
aagccccccg caaacagctg gccacgaaag ccgccaggaa aagcgctccc tctaccggcg	180
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<210> 71	
<211> 301	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1)...(301)	
<223> n = A,T,C or G	
<400> 71	
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ttgtttccctg cagccctccc acgggaatga caatggataa aagtgagctg gtacanaaaag	180
ccaaactcgc tgagcaggct gagcgatatg atgatatggc tgcagccatg aaggcagtca	240
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a	301
<210> 72	
<211> 251	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1)...(251)	
<223> n = A,T,C or G	
<400> 72	
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gtctctgtgc actctgtctt ggatgctctg gggagctcat gggtgagggaa gtctccacca	180
gagggaggct caggggactg gttggggcag ggatgaatat ttgaggggata aaaattgtgt	240
aagagccaan g	251
<210> 73	
<211> 895	
<212> DNA	
<213> Homo sapien	
<400> 73	
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caaattcttt ggtctcccat cagctgaat taatgtggta ctgtgtatct ttgagatcat	120
gtatttgtct ccactttgggt ggatacaaga aaggaaggca cgaacagctg aaaaagaagg	180
gtatcacacc gctccagctg gaatccagca ggaacctctg agcatgcac agctgaacac	240
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gtgcataaat ggtcatcata agtcaaacgt atcaattaga ccttcaacct aggaaaacaaa	360
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caaataaaat gcaggtgaaa gagatgaacc acgactagag gctgacttag aaatttatgc	540
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aagaattgtt aagaagtata ataacccttt caaaacccac aatgcagctt agtttcctt	720
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<210> 74	
<211> 351	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1)...(351)	
<223> n = A,T,C or G	
<400> 74	
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aggaaggagn catggcaagc catagctagg ccaccaatca gattaagaaa nnctgagaaaa	180
nctagctgac catcaactgtt ggtgnccagt ttcccaacac aatggaatnc caccacactg	240
gactagngga nccactagtt ctagagcggc cgccaccgcg gtggAACCCC aactttgcc	300
ccttagnga gggtaatttgc cgcgttggc ntaatcatgg tcataagctg t	351
<210> 75	
<211> 251	
<212> DNA	
<213> Homo sapien	
<400> 75	
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ccaaacataat tttttttaga tcgagtgcgc ataaatttct aagtcaGCCT ctatcggtgg	120
ttcatctctt tcacctgtcat ttatTTGGT gttgtctga agaaaggaaaa gagaaagca	180
aatacgaatt gtactatttgc taccaaatct ttggattca ttggcaataa atttcagtgt	240
ggtgttattat t	251
<210> 76	
<211> 251	
<212> DNA	
<213> Homo sapien	
<400> 76	
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tagtacaatt cgtatTTGCT ttccTCTTTC ctttCTTCAG acAAACACCA aataAAATGC	120
aggtgaaaga gatgaaccac gactagaggc tgacttagaa atttatGCTG actcgatcta	180
aaaaaaatTA TGTGGTTAA TGTAAATCTA TCTAAATAG AGCATTTGG gaatgcttt	240
caaagaaggt C	251
<210> 77	
<211> 351	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	

<222> (1)...(351)
 <223> n = A,T,C or G

<400> 77

actcaccgtg ctgtgtgctg tgtgcctgct gcctggcagc ctggccctgc cgctgctcag	60
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tatctctatg actcagaaac aaaaaatgcc aacagtttag aagccaaact caaggagatg	180
caaaaattct ttggcctacc tataactgga atgttaaact cccgcgtcat agaataatg	240
cagaagccca gatgtggagt gccagatgtt gcagaataact cactattcc aaatagccca	300
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<210> 78

<211> 1574

<212> DNA

<213> Homo sapien

<400> 78

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ccggcaacgg ctgcccgtc cctgggtccgg gcccaggggc ccgcgcctt ccgcggccgt	180
gctcgcgtg ctgctgtgc tcgccccgtt ggccggccccc gcgggggtccg gggaccccga	240
cgaccctggg cagcctcagg atgcgtgggtt cccgcgcagg ctccgcgcg aggccggcgc	300
cgccggcgctt cacttcttc acgtccgtc cgctcgcccc agcgcgcgtgc gagtgctggc	360
cgagggtgcag gagggccgcg cgtggattaa tccaaaagag ggtatgtaaag ttacacgttgt	420
cttcagcaca gagcgcctaca acccagagtc ttacttcag gaaggtgagg gacgtttggg	480
gaaatgttct gctcgagtgt tttaaagaa tcaaaaaaccc agaccaacta tcaatgtaac	540
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aatgaagcaa ctgaaaaacc ccttggaaat agtcagcata cctgataatc atggacatat	660
tgatccctct ctgagactca tctgggattt ggcttcctt ggaagctttt acgtgtatgt	720
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gaagtaccac tgtcaagagc tacagacacc agaagaagcc tccggaaactg aagaaggatc	960
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cattctaata atttggaaa acctatgtt acaagtaaaa actcagaaaat gcaaaagatgt	1140
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tttatgttgc taaaattgtt ctatctgttca tttttttttt aatggatgtt gctaaatata	1440
aagtaattt tagaaaaaaa attaactgtt ttaaaaagaa cttgattatg tttttttttt	1500
tcaggcaagt attcattttt aacttgcgttcc tttttttttt aatggatgtt gctaaatata	1560
aaaaaaaaaaaa aaaa	1574

<210> 79

<211> 401

<212> DNA

<213> Homo sapien

<220>

<221> misc_feature

<222> (1)...(401)

<223> n = A,T,C or G

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<400> 79
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ccagaaaacgt cacactgccc aagatggcca ggtacttcaa ggtctggAAC atgttgagct      180
gagtccagta gacatacatg agtcccagca tagcagcatg tcccaggtga aatataatcg      240
tgcttaggagc aaaagtgaag ttggagacat tggcaccaat ccggatccac tagttctaga      300
gcggccgcca ccgcgggtgga gctccagct ttgttccctt tagtgagggt taattgcgCG      360
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<210> 80
<211> 301
<212> DNA
<213> Homo sapien

<400> 80
aaaaatgaaa catctatTTT agcagcaaga ggctgtgagg gatggggtag aaaaggcatc      60
ctgagagagt tctagaccga cccaggtcct gtggcacact atacgggtca ggaggggtgg      120
aagacaggcc taagctctag gacggtaat ctccccctt tttgtggatt tgtagaaac      180
agacattctt ttggcctttt cctggcactg gtgtgcccgg caggtggca gaagtgagcc      240
accagtcaact gtcagtcat tgccaccaca gatcttcagc agaatcttcc ggtatcccc      300
t      301

<210> 81
<211> 301
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(301)
<223> n = A,T,C or G

<400> 81
tagccaggt gctcaagcta attttattct ttcccaacag gatccatttgc gaaaatatca      60
agcctttaga atgtggcagc aagagaaAGC ggactacgc ggaacgggga gtttgggaga      120
agctctcctg gtgttgactt agggatgaag gctccaggct gctgccagaa atggagtcac      180
cagcagaaga actgnTTCT ctgataagga tggccaccaca tttcaagct gttcgtaaa      240
tttacacagg tccttcttgc agcagtaagt accgttagct cattttccct caagcgggtt      300
t      301

<210> 82
<211> 201
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(201)
<223> n = A,T,C or G

<400> 82
tcaacagaca aaaaaagttt attgaataca aaactcaaAG gcatcaacAG tcctggggCC      60
aagagatCCA tggcaggaAG tcaagAGttc tgcttcaggG tcggctGGG cagccctGGA      120
agaagtcatG gcacatgaca gtgatgagtG ccaggAAAAC agcatactCC tggAAAGtCC      180
acctgctGGN cactgnTTCA t      201

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<210> 83
<211> 251
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(251)
<223> n = A,T,C or G

<400> 83

gtaaggagca tactgtgcc atttattata gaatgcagtt aaaaaaaaaata ttttgaggtt 60
agcctctcca gttaaaagg acttaacaag aaacacttgg acagcgatgc aatggtctct 120
cccaaaccgg ctcctctta ccaagtaccg taaacagggt ttgagaacgt tcaatcaatt 180
tcttgatatg aacaatcaa gcattaatg caaacatatt tgcttctcaa anaataaaaac 240
catttccaa a 251

<210> 84
<211> 301
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(301)
<223> n = A,T,C or G

<400> 84

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taaaaactca ggtatcagaa agactcaaaa ggctgtttt cactttgttc agattttgtt 120
tccaggcatt aagtgtgtca tacagtgtt gccactgctg tttccaaat gtccgatgtg 180
tgctatgact gacaactact tttctctggg tctgatcaat tttgcagttt accatttttag 240
ttcttacggc gtcnataaca aatgcttcaa catcatcagc tccaaatctga agtcttgctg 300
c 301

<210> 85
<211> 201
<212> DNA
<213> Homo sapien

<400> 85

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agtcacacca taaaggagtt tatccttaaa aggagtggaa gacattcaaa aaccaactgca 120
aataaaaaaaag ggtgacataa ttgctaaatg gagtggagga acagtgccta tcaattcttg 180
attgggccac aatgatatac c 201

<210> 86
<211> 301
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(301)

<223> n = A,T,C or G

<400> 86

tttataaaaat attttattta cagtagagct ttacaaaaat agtcttaaat taatacaat	60
ccctttgca atataactta tatgactatc ttctcaaaaa cgtgacattc gattataaca	120
cataaaactac atttatagtt gttaagtca cttgttagtat aaatatgttt tcatctttt	180
tttgtaataa ggtacatacc aataacaatg aacaatggac aacaaatctt atttgntat	240
tcttccaatg taaaattcat ctctggccaa aacaaaatta accaaagaaa agtaaaacaa	300
t	301

<210> 87

<211> 351

<212> DNA

<213> Homo sapien

<220>

<221> misc_feature

<222> (1)...(351)

<223> n = A,T,C or G

<400> 87

aaaaaaaaaggatt taagatcata aataggcat tgggtcaca acacattca gaatcttaaa	60
aaaaacaaaaca ttttggcttt ctaagaaaaa gactttaaa aaaaatcaat tccctcatca	120
ctgaaaggac ttgtacattt ttaaacttcc agtctcctaa ggcacagtagt ttaatcagaa	180
tgc当地atatt accaccctgc tggc当地anga ataaagaagc aagggatata cactaaaaa	240
aacngccaaa ttcttgaacc aaatcattgg catttaaaa aagggataaa aaaacnggnt	300
aagggggggga gcatttttaag taaagaang ccaagggtgg tatgccnnga c	351

<210> 88

<211> 301

<212> DNA

<213> Homo sapien

<220>

<221> misc_feature

<222> (1)...(301)

<223> n = A,T,C or G

<400> 88

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ttttctggtg gaagcacaca gtaatttac tcaagtgtgg cgntagcgat gctttttcat	180
ggngtcatattt atccacttgg tgaacttgca cacttgaatg naaactcctg ggtcattggg	240
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c	301

<210> 89

<211> 591

<212> DNA

<213> Homo sapien

<400> 89

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aagcaccagg ctggtcctct ccccacatgt cacactctcc tcagcctctc ccccaaccct	120
gctctccctc ctccccctgcc ctagccagg gacagagtct aggaggagcc tggggcagag	180

ctggaggcag	gaagagagca	ctggacagac	agctatggtt	tggattgggg	aagagattag	240
gaagtaggtt	cttaaagacc	cttttttagt	accagatatac	cagccatatt	cccagctcca	300
ttattcaaat	catttccat	agcccagctc	ctctctgttc	tccccctact	accaattctt	360
tggctcttac	acaatttta	tccctcaaat	attcatccct	ggcccaacca	gtcccctgag	420
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acagagacct	gggaaaggaa	gctgaactt	gcagagatgt	ggacagggtgc	aggctagggt	540
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<210> 90
<211> 1978
<212> DNA
<213> Homo sapien

<400> 90

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caagatctaa	cagggttctt	tctagaggat	tattagataa	gtaacacttg	atcattaagc	660
acggatcatg	ccactcattc	atgggttcttc	tatgttccat	gaactcta	atgcccac	720
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ggtgactcag	caaccaaaca	aaaggata	taattttta	tgaacaat	atttgtattt	1920
tatggacata	aaaggaaact	ttcagaaaga	aaaggaggaa	aataaagggg	gaaaggga	1978

<210> 91
<211> 895
<212> DNA
<213> Homo sapien

<400> 91

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aaacgctgga	ctacaaaatgc	aggtttcttt	atatccttaa	cttcaattat	tgtcacttat	180
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taagttaatg	cacaaccaag	agtatacact	gttcatttgt	gcagttatgc	gtcaaatgcg	300
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atagattgaa	atacagtttA	aaacaaaat	tgtattctac	aaatacaata	aaatttgcAA	420
cttgcacatc	tgaagcaaca	tttgagaaag	ctgcttcaat	accCCTGCTG	ttatattggT	480
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atgcatgagt	ccatctacat	atTTTCAAGT	tttggaaaca	gaaagaAGTT	tagaattttc	660
aaagtaatct	gaaaACTTTC	taagccattt	taaaataaga	tttttttccc	catctttcca	720
atgtttccta	tttgatagtgt	taatacagaa	atgggcagtt	tctagtgtca	acttaactgt	780
gctaattcat	aagtcaTTAT	acatttATGA	cttaagAGTT	caaataAGTG	gaaattgggt	840
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<210> 92
<211> 1692
<212> DNA
<213> Homo sapien

<400> 92						
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ttacaaAGCA	tcttatttGAT	ttaaaAGAT	ccatactatt	gataaAGTC	accatgaACA	180
tatATGTAAT	aaggagactA	aaatattCAT	tttacatATC	tacaACATGT	atTTCAATT	240
tctaATCAAC	cacaATCAT	ataggAAAAT	atTTAGGTCC	atgAAAAGT	ttcaAAACAT	300
taaaaaATTAA	aagTTTgAA	acaatTCACA	tgtgAAAGCT	cattaaATAA	taacATTGAC	360
aaATAAATAG	ttaATCAGCT	ttacttATTAA	gctgctGCCA	tgcatttCTG	gcattCCATT	420
ccaAGCGAGG	gtcagcatGC	agggtataAT	ttcataCTAT	gcgaccGTA	agagCTACAG	480
ggCTTATTtT	tgaagtgAAA	tgtcacAGGG	tcttcatTC	tcttcaAG	gaagatCACT	540
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<213> Homo sapien

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<212> DNA
<213> Homo sapien

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<212> DNA
<213> Homo sapien

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<211> 594
<212> DNA
<213> Homo sapien

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<212> DNA
<213> Homo sapien

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<210> 98
<211> 90
<212> PRT
<213> Homo sapien

<400> 98
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35 40 45
Ala Ala Thr Thr Ala Thr Thr Ala Ala Pro Thr Thr Ala Thr Thr Ala
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Ala Ser Thr Thr Ala Arg Lys Asp Ile Pro Val Leu Pro Lys Trp Val
65 70 75 80
Gly Asp Leu Pro Asn Gly Arg Val Cys Pro
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<210> 99
<211> 197
<212> PRT
<213> Homo sapien

<400> 99
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35 40 45
Glu Lys Leu Trp Thr Glu Val Asn Ala Leu Lys Glu Ile Gln Ala Leu
50 55 60
Gln Thr Val Cys Leu Arg Gly Thr Lys Val His Lys Lys Cys Tyr Leu
65 70 75 80
Ala Ser Glu Gly Leu Lys His Phe His Glu Ala Asn Glu Asp Cys Ile
85 90 95
Ser Lys Gly Gly Ile Leu Val Ile Pro Arg Asn Ser Asp Glu Ile Asn
100 105 110
Ala Leu Gln Asp Tyr Gly Lys Arg Ser Leu Pro Gly Val Asn Asp Phe
115 120 125
Trp Leu Gly Ile Asn Asp Met Val Thr Glu Gly Lys Phe Val Asp Val
130 135 140
Asn Gly Ile Ala Ile Ser Phe Leu Asn Trp Asp Arg Ala Gln Pro Asn

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		Gln	Gly
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<211> 3410
<212> DNA
<213> Homo sapien

<400> 100

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<210> 101
<211> 553
<212> PRT
<213> Homo sapien

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Gly Thr Gln Glu Glu Cys Leu Phe Gly Leu Leu Thr Leu Ile Phe Leu	
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Pro Thr Glu Pro Ala Glu Gly Leu Ser Ala Pro Ser Leu Ser Pro His	
225 230 235 240	
Cys Cys Pro Cys Arg Ala Arg Leu Ala Phe Arg Asn Leu Gly Ala Leu	
245 250 255	

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 355 360 365
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 Gly Gly Ser Gly Leu Leu Pro Pro Pro Pro Ala Leu Cys Gly Ala Ser
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 Arg Val Val Pro Gly Arg Gly Ile Cys Leu Asp Leu Ala Ile Leu Asp
 485 490 495
 Ser Ala Phe Leu Leu Ser Gln Val Ala Pro Ser Leu Phe Met Gly Ser
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 Ile Val Gln Leu Ser Gln Ser Val Thr Ala Tyr Met Val Ser Ala Ala
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 Lys Ser Asp Leu Ala Lys Tyr Ser Ala
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<210> 102

<211> 940

<212> DNA

<213> Human

<400> 102

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